

(FILE 'HOME' ENTERED AT 15:34:50 ON 12 JUL 2000)

FILE 'BIOSIS, MEDLINE, CAPLUS, EMBASE, SCISEARCH' ENTERED AT 15:35:51 ON 12 JUL 2000

	12 JUL 200	U Company of the comp
L1	6	S JOINT TIME FREQUENCY TRANSFORM
L2	4	DUPLICATE REMOVE L1 (2 DUPLICATES REMOVED)
L3	7155	S FAST FOURIER TRANSFORM
L4	3305826	S NUCLEIC ACID? OR OLIGONUCLEOTIDE? OR DNA OR RNA
L5	56	S L3 AND L4
L6	193452	S ARRAY?
L7	576687	S ELECTRODE
L8	0	S L5 AND L6 AND L7
L9	1	S L5 AND L6
L10	7127	S ALTERNATING CURRENT?
L11	2606	S HARMONIC ANALYSIS
L12	0	S L4 AND L10 AND L11
L13	87	S L4 AND L10
L14	21	S L4 AND L11
L15	0	S L14 AND L6 AND L7
L16	0	S L14 AND L7
L17	0	S L14 AND L6
L18	8	DUPLICATE REMOVE L14 (13 DUPLICATES REMOVED)
L19	0	S L4 AND L6 AND L7 AND L10
L20	0	S L4 AND L6 AND L10
L21	38	S L4 AND L7 AND L10
L22	23	DUPLICATE REMOVE L21 (15 DUPLICATES REMOVED)
L23		S PEAK RECOGNITION
L24	3	S L4 AND L23
L25		S PROCESSING
L26	17	S L23 AND L25
L27	14	DUPLICATE REMOVE L26 (3 DUPLICATES REMOVED)
L28	0	S L10 AND L23
L29	6249	S DIGITAL FILTER?
L30		S L4 AND L29
L31		S L7 AND L30
L32	0	
L33	2872	S SIGNAL AVERAGING
L34		S L10 AND L33
L35		DUPLICATE REMOVE L34 (6 DUPLICATES REMOVED)
L36	1	
L37	_	S SPECTRAL ANALYSIS
L38		S L10 AND L37
L39	0	S L4 AND L38
L40	•	S L7 AND L38
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ANSWER 1 OF 4 SCISEARCH COPYRIGHT 2000 ISI (R) 1998:310007 SCISEARCH AN The Genuine Article (R) Number: ZH484 GΑ Joint time-frequency transform for TIradar range Doppler imaging Chen V C (Reprint); Qian S ΑU USN, RES LAB, DIV RADAR, 4555 OVERLOOK AVE SW, WASHINGTON, DC 20375 CS (Reprint); NATL INSTRUMENTS CORP, DSP GRP, AUSTIN, TX 78730 CYA USA SO IEEE TRANSACTIONS ON AEROSPACE AND ELECTRONIC SYSTEMS, (APR 1998) Vol. 34, No. 2, pp. 486-499. Publisher: IEEE-INST ELECTRICAL ELECTRONICS ENGINEERS INC, 345 E 47TH ST, NEW YORK, NY 10017-2394. ISSN: 0018-9251. DTArticle; Journal FS ENGI English LAREC Reference Count: 17 Conventional radar imaging uses the Fourier transform to retrieve AB Doppler information. However, due to the complex motion of a target, the Doppler frequency shifts are actually time-varying. By using the Fourier transform, the Doppler spectrum becomes smeared and the image is blurred. Without resorting to sophisticated motion compensation algorithms, the image blurring problem can be resolved with the joint time-frequency transform. High-resolution time-frequency transforms are investigated, and examples of applications to radar imaging of single and multiple targets with complex motion are given. AEROSPACE ENGINEERING & TECHNOLOGY; ENGINEERING, ELECTRICAL & ELECTRONIC CCKeyWords Plus (R): WIGNER DISTRIBUTION; SIGNAL ANALYSIS; TOOL STP RE |Year | VOL | PG | Referenced Work Referenced Author |(RPY)|(RVL)|(RPG)| (RWK) (RAU) ______+ |1984 |20 | 363 | IEEE T AERO ELEC SYS AUSHERMAN D A |1995 | |CH4 |SPOTLIGHT SYNTHETIC CARRARA W G |2212 | OPT ENG |1994 |33 CHEN V C CHEN V C CLAASEN T A |1067 | PHILLIPS J RES |1980 |35 |217 | PHILIPS J RES |1980 |35 CLAASEN T A C M |276 | PHILIPS J RES CLAASEN T A C M |1980 |35

T.2	ANSWER	2	OF	4	MEDLINE

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WOODWARD P M

WEHNER D R

DAVENPORT W B

TI Time-frequency transforms: a new approach to first heart sound frequency dynamics.

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AU Wood J C; Buda A J; Barry D T

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|SIGNAL PROCESS

|CH6 | HIGH RESOLUTION RADA

|CH7 | PROBABILITY INFORMAT

Department of Internal Medicine, University of Michigan Medical School, CS Ann Arbor 48109... NS01701 (NINDS) NC HL34691 (NHLBI) IEEE TRANSACTIONS ON BIOMEDICAL ENGINEERING, (1992 Jul) 39 (7) 730-40. SO Journal code: GFX. ISSN: 0018-9294. CYUnited States Journal; Article; (JOURNAL ARTICLE) \mathtt{DT} $\mathtt{L}\mathtt{A}$ English 199212 EMThis study employed a new analytical tool, the Binomial joint AB time-frequency transform, to test the hypothesis that first heart sound frequency rises during the isovolumic contraction period. Cardiac vibrations were recorded from eight open chest dogs using an ultralight accelerometer cemented directly to the epicardium of the anterior left ventricle. The frequency response of the recording system was flat +/- 3 dB from 0.1 to 400 Hz. Three characteristic time-frequency spectral patterns were evident in the animals investigated: 1) A frequency component that rose from approximately 40-140 Hz in a 30-50 ms interval immediately following the ECG R-wave. 2) A slowly varying or static frequency of 60-100 Hz beginning midway through the isovolumic contraction period. 3) Broad-band peaks occurring at the time of the Ia and Ib high frequency components. The presence of rapid frequency dynamics limits the usefulness of stationary analysis techniques for the first heart sound. The Binomial transform provided much better resolution than the spectrograph or spectrogram, the two most common non-stationary signal analysis techniques. By revealing the onset and dynamics of first heart sound frequencies, time-frequency transforms may allow mechanical assessment of individual cardiac structures. Check Tags: Animal; Male; Support, Non-U.S. Gov't; Support, U.S. Gov't, CTNon-P.H.S.; Support, U.S. Gov't, P.H.S. Dogs Evaluation Studies Fourier Analysis *Heart Sounds *Hemodynamics Phonocardiography: MT, methods *Phonocardiography: ST, standards *Signal Processing, Computer-Assisted ANSWER 3 OF 4 SCISEARCH COPYRIGHT 2000 ISI (R) L2AN92:366761 SCISEARCH The Genuine Article (R) Number: HY634 GΑ WIGNER DISTRIBUTION DECOMPOSITION AND CROSS-TERMS DELETED REPRESENTATION TIΑU SHIE Q (Reprint); MORRIS J M NATL INSTRUMENTS, DIV DSP, 6504 BRIDGE POINT PKWY, AUSTIN, TX, 78730 CS (Reprint) ÇYA USA SIGNAL PROCESSING, (MAY 1992) Vol. 27, No. 2, pp. 125-144. SO ISSN: 0165-1684. Article; Journal DTFS ENGI ENGLISH LA REC No References Keyed In this paper, we represent the Wigner Distribution (WD) of an AB arbitrary signal, via the Gabor expansion, in terms of a linear

combination of elementary WDs, which can be easily partitioned into two subsets: auto WDs and cross WDs. The Gabor coefficients, C(m,n) for this

decomposition are obtained with a Gaussian-shaped synthesis function. The optimally concentrated auto WDs are non-negative and ntirely free of cross-terms; the sum of these auto WDs we call the constraint deleted representation (CDR). The sum of the cross WDs is an oscillating function with non-zero energy in general; it can be removed and returned depending on the user's needs. Such a decomposition illustrates and isolates the mechanism of WD negative values and cross-term interference. Moreover,

new

information is provided to facilitate the design of valid joint time-frequency signal representations and time-varying filters. Also in this paper, analogous, yet more practical, results are shown for the Discrete Wigner Distribution (DWD) for finite or periodic discrete-time signals. Examples are presented to demonstrate the CDR technique and its performance in comparison with other joint time-frequency distributions. It is shown that the CDR has the high energy concentration of the WD without the interference problems that occur in many other approaches. Moreover, because only the Gabor coefficients, C(m,n), need be computed on-line, the CDR is suitable for on-line implementation.

- CC ENGINEERING, ELECTRICAL & ELECTRONIC
- Author Keywords: WIGNER DISTRIBUTION; GABOR EXPANSION; JOINT

 TIME FREQUENCY TRANSFORMS; CROSS-TERM

 INTERFERENCE; SPECTROGRAM; CHOI-WILLIAMS DISTRIBUTION; DISCRETE WIGNER

 DISTRIBUTION
- L2 ANSWER 4 OF 4 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1991:39582 BIOSIS
- DN BR40:16562
- TI NEW EVIDENCE FOR MYOCARDIAL GENESIS OF THE FIRST HEART SOUND.
- AU WOOD J C; BARRY D T; GALLAGHER M; HAARER S; BUDA A J
- CS UNIV. MICH. MED. SCH., ANN ARBOR, MICH.
- SO 63RD SCIENTIFIC SESSIONS OF THE AMERICAN HEART ASSOCIATION, DALLAS, TEXAS,

USA, NOVEMBER 12-15, 1990. CIRCULATION. (1990) 82 (4 SUPPL 3), III578. CODEN: CIRCAZ. ISSN: 0009-7322.

- DT Conference
- FS BR; OLD
- LA English
- CC General Biology Symposia, Transactions and Proceedings of Conferences, Congresses, Review Annuals 00520 Cardiovascular System General; Methods *14501 Cardiovascular System Physiology and Biochemistry *14504
- BC Canidae 85765
- IT Miscellaneous Descriptors

ABSTRACT DOG JOINT TIME FREQUENCY

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9 ANSWER 1 OF 1 BIOSIS COPYRIGHT 2000 BIOSIS
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AN 1991:268013 BIOSIS

DN BA92:628

TI FAST FOURIER TRANSFORM-BASED CORRELATION OF DNA SEQUENCES USING COMPLEX PLANE ENCODING.

AU CHEEVER E A; OVERTON G C; SEARLS D B

CS CENT. ADVANCED INFORMATION TECHNOL., UNISYS CORP., P.O. BOX 517, PAOLI, PA. 19301.

SO COMPUT APPL BIOSCI, (1991) 7 (2), 143-154. CODEN: COABER. ISSN: 0266-7061.

FS BA; OLD

LA English

AB The detection of similarities between DNA sequences can be accomplished using the signal-processing technique of cross-correlation. An early method used the fast Fourier

transform (FFT) to perform correlations on DNA sequences in O(n log n) time for any length sequence. However, this method requires many FFTs (nine), runs no faster if one sequence is much shorter than the other, and measures only global similarity, so that significant short local matches may be missed. We report that, through the use of alternative encodings of the DNA sequence in the complex plane, the number of FFTs performed can be traded off against (1)

signal-to-noise

ratio, and (ii) a certain degree of filtering for local similarity via k-tuple correlation. Also, when comparing probe sequences against much longer targets, the algorithm can be sped up by decomposing the target

and

performing multiple small FFTs in an overlap-save arrangment. Finally, by decomposing the probe sequence as well, the detection of local similarities can be further enhanced. With current advances in extremely fast hardware implementations of signal-processing operations, this approach may prove more practical than heretofore.

General Biology - Information, Documentation, Retrieval and Computer Applications *00530

Mathematical Biology and Statistical Methods *04500

Biochem

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ANSWER 1 OF 8 BIOSIS COPYRIGHT 2000 BIOSIS
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       2000:253260 BIOSIS
  AN
       PREV200000253260
  DN
       Harmonic analysis of DNA dynamics in a
  TI
       viscous medium.
       Shih, Chia C. (1); Georghiou, S. (1)
  ΑU
       (1) Department of Physics, University of Tennessee, Knoxville, TN,
  CS
       37996-1200 USA
       Journal of Biomolecular Structure and Dynamics, (April, 2000) Vol. 17,
  SO
 No.
       5, pp. 921-932. print..
      ISSN: 0739-1102.
 DT
      Article
      English
 LA
      English
 SL
      The harmonic dynamics of normal modes of double-stranded DNA in
 AB
      a viscous fluid are investigated. The model DNA consists of two
      backbone-supported DNA strands coiling around a common helix
      axis with base stacking, sugar puckering, interstrand hydrogen bonding,
      and intrastrand sugar-base interactions assigned values based on
 published
      data. Assuming that the DNA bases are shielded from direct
      bombardment by the solvent, analytical solutions are obtained. The
      dissipation and fluctuation of the normal modes of the bases moving along
      the spirals display the effect of the medium indirectly through
      interactions with the backbone. The dynamics of the backbone are found to
      be overdamped with the characteristic damping times extending to the
      picosecond region for disturbance in position and to the sub-picosecond
      region for disturbance in velocity. In addition to the dynamic mode of a
      rigid rod, the motions of the bases are coupled to the motions of the
      backbone with comparable amplitudes for disturbance in position. For
      disturbance in velocity, however, the bases are effectively at rest, not
      being able to follow the motions of the backbone. The angular frequencies
      of the underdamped vibrational modes, identified as the ringing modes of
      the bases with the backbone effectively at rest, are insensitive to the
      viscosity and lie in the low frequency region of the Raman spectrum.
These -
      findings indicate that the backbone of DNA plays a significant
      role in modulating the dynamics of double-stranded DNA in an
     overdamping environment. This modulation of the dynamics of the motions
ο£
     the bases in DNA by environmental impediments to molecular
     motion is briefly discussed in connection with protein- and drug-
     DNA interactions as well as gene regulation.
     Biochemical Studies - Nucleic Acids, Purines and Pyrimidines *10062
CC
     Biophysics - Biocybernetics *10515
     Major Concepts
IT
        Biochemistry and Molecular Biophysics; Models and Simulations
        (Computational Biology)
     Chemicals & Biochemicals
ΙŢ
        double-stranded DNA
     Miscellaneous Descriptors
ΙT
        DNA dynamics: harmonic analysis; drug-
      DNA interactions; gene regulation; molecular motion; protein-
      DNA interaction; viscous medium
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    1998:548964 CAPLUS
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    129:286999
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Art Unit: 1655

It would have been of Il in the art at the time the invention was made to have synthesized more sources of mRNA using a primer containing a specific se are a T7 promoter sequence as suggested by Logel et al. and have prod c' A from a cDNA library representing two or more sources of mRN clabeled primer as suggested by Luehrsen et al. and T7 polyme as to et al.. The prior arts provided by Sagerstrom et al., Loge et al as motivated one having ordinary skill in the art to test the pos NAs from two or more sources of mRNA using a primer contain ete I from T3 or T7 promoter sequence and produce differen bury representing two or more sources of mRNA in the preis suggested by Luehrsen et al. and T7 polymerase. One have e time the invention was made would have been a reasonable expect and ads together because all of these methods are known in '

- 11. No claim is allowed.
- 12. Papers related to thi.

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(December 28, 1993 (See 37)

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W. Gary Jones, can be reached

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Frank Lu July 12, 2000 1. : 16, 1993), and 1157 OG 94

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14. the examiner's supervisor,

s application should be

is (703) 308-0196.

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Continuum Solvent Studies of the Stability of DNA, RNA
     Srinivasan, Jayash :; Cheatham, Thomas E., III; Cie, Lak, Piotr; Kollman,
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ΑU
     Department of Molecular Biology, Scripps Research Institute, La Jolla,
ÇS
CA,
     92037, USA
     J. Am. Chem. Soc. (1998), 120(37), 9401-9409
SO
     CODEN: JACSAT; ISSN: 0002-7863
     American Chemical Society
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     We apply continuum solvent models to investigate the relative stability
CC
AΒ
     A- and B-form helixes for three DNA sequences, d(CCAACGTTGG)2,
οf
      d(ACCCGCGGGT)2, and d(CGCGAATTCGCG)2, a phosphoramidate-modified
      DNA duplex, p(CGCGAATTCGCG)2, in which the O3' atom in deoxyribose
      is replaced with NH, and an RNA duplex, r(CCAACGUUGG) 2.
      Structures were taken as snapshots from multi-nanosecond mol. dynamics
      simulations computed in a consistent fashion using explicit solvent and
      with long-range electrostatics accounted for using the particle-mesh
      procedure. The electrostatic contribution to solvation energies were
      computed using both a finite-difference Poisson-Boltzmann (PB) model and
 Ewald
      pairwise generalized Born model; nonelectrostatic contributions were
 a
      with a surface-area-dependent term. To these solvation free energies
  estd.
       added the mean solute internal energies (detd. from a mol. mechanics
  were
       potential) and ests. of the solute entropy (from a harmonic
       anal.). Consistent with expt., the relative energies favor B-form
       helixes for DNA and A-form helixes for the NP-modified system
       and for RNA. Salt effects, modeled at the linear or nonlinear
       PB level, favor the A-form helixes by modest amts.; for d(ACCCGCGGGT)2,
       salt is nearly able to switch the conformational preference to "A". The
       results provide a phys. interpretation for the origins of the relative
       stabilities of A- and B-helixes and suggest that similar analyses might
       useful in a variety of nucleic acid conformational
  be
       problems.
       DNA RNA helix conformation stability
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        RNA, and phosphoramidate-DNA helixes)
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m IT}
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           (continuum solvent studies of the stability of DNA,
         RNA, and phosphoramidate-DNA helixes)
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        RL: PRP (Properties)
           (continuum solvent studies of the stability of DNA,
         RNA, and phosphoramidate-DNA helixes)
        ANSWER 3 OF 8 SCISEARCH COPYRIGHT 2000 ISI (R)
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        1998:323687 SCISEARCH
        The Genuine Article (R) Number: ZJ230
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        Microfibrillar buckling within fibers under compression
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         JOURNAL OF CHEMICAL PHYSICS, (22 APR 1998) Vol. 108, No. 16, pp.
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Art Unit: 1655

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The teaching of Sag and Sagerstrom and I al. do recommend to the company of the sagerstrom and Topolymerase.

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el et al., (Biotechnique 13, 604-17), and Bodescot et al., (DNA Cell

fter subtracting 1st strand cDNA ver), a tag sequence selected from resence of fluorescence labeled

Teviously, supra.

A cDNA (tracer) from tissue A with trary and synthesis of cDNA in the

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of previously, supra.

Old cDNA (tracer) from tissue A

ONA in the presence of T7

nt' osis using T7 DNA polymerase

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    BLVD, WOODBURY, NY, 797-2999.
    ISSN: 0021-9606.
    Article; Journal
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FS
    English
        A tentative theory is presented of microfibrillar buckling within
LA
    Reference Count: 22
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     compressed fibers. A quantitative harmonic analysis is
     given of the semiclassical buckling of a clamped stiff chain; the
AB
     influence of thermal undulations is incorporated in Euler buckling. A
     scaling analysis including entropy allows one to understand semiclassical
     buckling. The buckling of a microfibril within a fibrous environment is
     analyzed in two limits: (a) when the fiber is incompressible; (b) when
     matrix is assumed to be a fixed harmonic potential. In the latter case, a
     network of microfibrils may melt at high enough compression before the
 the
     usual bucking occurs. We also study the renormalization of the confining
     potential by long-range elastic fields. A provisional comparison with
     experimental studies on macroscopic failure is given. (C) 1998 American
      Institute of Physics.
      KeyWords Plus (R): WORMLIKE CHAINS; POLYMERS; COMPOSITES; FILAMENTS;
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      STRENGTH; MODEL; DNA
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         Continuum solvent studies of the stability of RNA hairpin loops
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         Srinivasan, Jayashree; Miller, Jennifer; Kollman, Peter A.; Case, David
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    AU
         (1) Dep. Molecular Biol., Scripps Res. Inst., La Jolla, CA 92037 USA
    Α.
         Journal of Biomolecular Structure and Dynamics, (Dec., 1998) Vol. 16, No.
    CS
     SO
          3, pp. 671-682.
          ISSN: 0739-1102.
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Application/Control Number: (19/31 09

Art Unit: 1655

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Jescript II SK (+/-) phagemid vector e. T7 and T3 promoter sequences. kill in the art at the time the invention or more sources of mRNA using a ication, a selected marker gene, a T7 ordonuclease site distal to the tailed atagene Catalogue, and have e sources of mRNA by combining 1st A commercially available pBluescript II perstrom et al. and Belyavsky et al. to test the possibility to synthesize 1st hemi-tailed vector or primer T7 or T3 promoter sequence and at ed terminus of the plasmid and ml NA together in order to produce a NA. One having ordinary skill in the sonable expectation of success to he stare known in the art and are easy

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English
  LA
       We apply continuum olvent models to investigate the relative stability
  AB
  οf
       various conformational forms for two RNA sequences,
       GGAC (UUCG) GUCC and GGUG (UGAA) CACC. In the first part, we compare
  alternate
       hairpin conformations to explore the reliability of these models to
       discriminate between different local conformations. A second part looks
  at
       the hairpin-duplex conversion for the UUCG sequence, identifying major
      contributors to the thermodynamics of a much large scale transition.
      Structures were taken as snapshots from multi-nanosecond molecular
      dynamics simulations computed in a consistent fashion using explicit
      solvent and with long-range electrostatics accounted for using the
      Particle-Mesh Ewald procedure. The electrostatic contribution to
 solvation
      energies were computed using both a finite-difference Poisson-Boltzmann
      (PB) model and a pairwise Generalized Born model; non-electrostatic
      contributions were estimated with a surface-area dependent term. To these
      solvation free energies were added the mean solute internal energies
      (determined from a molecular mechanics potential) and estimates of the
      solute entropy (from a harmonic analysis). Consistent
      with experiment and with earlier solvated molecular dynamics simulations,
      the UUCG hairpin was found to prefer conformers close to a recent NMR
      structure determination in preference to those from an earlier NMR study.
      Similarly, results for the UGAA hairpin favored an NMR-derived structure
      over that to be expected for a generic GNRA hairpin loop. Experimental
      free energies are not known for the hairpin/duplex conversion, but must
 1717
      close to zero since hairpins are seen in solution and duplexes in
      crystals; out calculations find a value near zero and illustrate the
      expected interplay of solvation, salt effects and entropy in affecting
      this equilibrium.
      Brochemical Studies - Nucleic Acids, Purines and Pyrimidines *10062
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      Genetics and Cytogenetics - General *03502
      Comparative Biochemistry, General *10010
      Biochemical Methods - Nucleic Acids, Purines and Pyrimidines *10052
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     Biophysics - Molecular Properties and Macromolecules *10506
     Major Concepts
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         Biochemistry and Molecular Biophysics
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        solvents; RNA: hairpin loops, helices, molecular
        characteristics, stabilities
     Methods & Equipment
IT
        NMR spectroscopy: analytical method, spectroscopic techniques: CB
     Miscellaneous Descriptors
IT
        continuum solvent studies; electrostatics; free energy; thermodynamics
L18 ANSWER 5 OF 8 SCISEARCH COPYRIGHT 2000 ISI (R)
     94:649709 SCISEARCH
AN
     The Genuine Article (R) Number: PK386
GA
     EMERGENCE OF REGULAR SUPERSTRUCTURES IN MACROMOLECULES
TI
     BOTTI S A (Reprint); DESANTIS P; FUA M
AU
     UNIV ROMA LA SAPIENZA, DIPARTIMENTO CHIM, I-00185 ROME, ITALY (Reprint)
CS
CYA ITALY
     BERICHTE DER BUNSEN GESELLSCHAFT FUR PHYSIKALISCHE CHEMIE-AN
SO
INTERNATIONAL
     JOURNAL OF PHYSICAL CHEMISTRY, (SEP 1994) Vol. 98, No. 9, pp. 1194-1197.
     ISSN: 0005-9021.
    Article; Journal
\mathsf{DT}
     PHYS
FS
LA
    ENGLISH
REC Reference Count: 8
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the second of th

Application/Control Number: 09/350,609

Art Unit: 1655

9. Claims 13 is rejected under 35 U.S.C. 103(
al., (Annu. Rev. Biochem. 66, 751-783, 1997) in virialised on July 11, 1995), Margolskee at 'S Patent 5,2
Stratagene Catalogue (1994, page 298).

The teachings of Sager from et al. have be-

Sagerstrom et al. do not discome cDNA construction (tracer) from tissue A with poly(A). RNA from tisprimer comprising an origin of a olication, a selecter and at least one unique restriction en fonuclease site.

The teachings of Belyavsky ///. have bee

Belyavsky *et al.* do not sclose the subtract with poly(A)+RNA from tissue Bassiver) and a horizon of replication, a selected management, a T7 unique restriction endonuclease life distal to the talent school of the subtraction of the subtrac

Margolskee teach high-citienty cloning et (see Figure 1).

Margolskee does not dis dos. DNA cons (tracer) from tissue A with 1 d. A. TNA from the primer comprising a T7 or T3 1 or at sequence.

Eing unpatentable over Sagerstrom et Edyavsky et al., (US Patent 5,814, 445, filed on August 12, 1992), and

invarized previously, supra.

ion after subtracting 1st strand cDNA

(criver) and a hemi-tailed vector or

ker gene, a T7 or T3 promoter sequence

to the tailed terminus of the plasmid.

arrized previously, supra.

Ist strand cDNA (tracer) from tissue A like I vector or primer comprising an moter sequence and at least one minus of the plasmid.

using a hemi-tailed vector or primer

v subtracting 1st strand cDNA
viver) and a hemi-tailed vector or

AB A general method is described in which the harmonic analysis of perturbations is applied to the study of superstructures of cromolecular chains. The theoretical approach employed has been to apply harmonic perturbations on the conformational parameters in macromolecular helices of various periodicities and to study the overall variation in structure and its dependency on the periodicity

the overall variation in structure and its dependency on the periodicity of the perturbation. The results clearly show that when these perturbations do not contain harmonics close to the fundamental periodicities of the polymer chain, the consequent structural effects remain localized and are not productive at a superstructural level. Furthermore, the features of these superstructures are dependent only on the amplitude of the fundamental periodicity component of the perturbation

and are generated by topologically equivalent transformations. These findings enable us to devise a model to study and identify transconformational pathways leading to global variations in the structure

of the macromolecular chain.

CC CHEMISTRY, PHYSICAL

Author Keywords: BIOLOGICAL MACROMOLECULES; COMPUTER EXPERIMENTS; POLYMER STP KeyWords Plus (R): THEORETICAL PREDICTION; CURVATURE; SEQUENCE; PROTEINS; DNAS

RE

Referenced Author (RAU)	Year VOL PG (RPY) (RVL) (RPG	, , , , , , , , , , , , , , , , , , , ,
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		PROTEINS
BOFFELLI D	1991 39 127	BIOPHYS CHEM
BOFFELLI D	1992 42 1409	IINT J QUANTUM CHEM
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DESANTIS P	1984 23 1547	BIOPOLYMERS
DESANTIS P	1985	STRUCTURE MOTION MEM
MOROSETTI S	1974 7 52	MACROMOLECULES
TRIFONOV E N	1980 77 13816	IP NATL ACAD SCI USA

- L18 ANSWER 6 OF 8 SCISEARCH COPYRIGHT 2000 ISI (R)
- AN 94:15691 SCISEARCH
- GA The Genuine Article (R) Number: MM819
- TI SPECTRAL-ANALYSIS FOR CATEGORICAL TIME-SERIES SCALING AND THE SPECTRAL ENVELOPE
- AU STOFFER D S (Reprint); TYLER D E; MCDOUGALL A J
- CS UNIV PITTSBURGH, DEPT MATH & STAT, PITTSBURGH, PA, 15260 (Reprint); RUTGERS UNIV, DEPT STAT, NEW BRUNSWICK, NJ, 08903

CYA USA

- SO BIOMETRIKA, (SEP 1993) Vol. 80, No. 3, pp. 611-622. ISSN: 0006-3444.
- DT Article; Journal
- FS PHYS; LIFE; AGRI
- LA ENGLISH

AB

REC Reference Count: 22

Many studies produce categorical time series in which harmonic analysis is of interest. Although there exist time domain approaches for the analysis of categorical time series such as Markov chains or link function based regression models, there is apparently little statistical theory or methodology for analyzing qualitative-valued time series in the frequency domain. The purpose of this paper is to initiate the development of a general framework for the frequency domain analysis of categorical time series. In doing so, we discuss the scaling of categorical time series and introduce a new concept that we call the spectral envelope of a categorical time series. We demonstrate our methodology on a data set from a problem in molecular biology.

CC MATHEMATICAL METHODS, BIOLOGY & MEDICINE; STATISTICS & PROBABILITY

Author Keywords: ASYMPTOTIC DISTRIBUTION OF LATENT ROOTS AND VECTORS; DNA SEQUENCING; FREQUENCY DOMAIN ANALYSIS; MARKOV CHAIN;

Application/Control Number: 09/350,609

Art Unit: 1655

Logel et al. do not disclose the subtraction of 1:1 stand cDNA (tracer) from tissue A with poly(A)+ RNA from tissue B (driver) and production if cDNA library.

Luehrsen et al., teach analysis of differential d primers and genescanTM software(page 168, abstract) presence of fluorescence labeled poly(dT) (page 170)

It would have been obvious to one having ord was made to have synthesized 1st strand cDNAs from hemi-tailed primer containing a specific sequence tages sequence as suggested by Logel et al. and have proor more sources of mRNA by combining 1st strand cl al. and synthesizing 2nd strand cDNA in the presence dG oligonucleotide tail to the 3' termini of the hetere et al.. The prior arts provided by Sagerstrom et al. one having ordinary skill in the crt to test the possil iii or more sources of mRNA using a hemi-tailed prim from T3, T7, and SP6 promoter sequence and comisources of mRNA together in order to produce a sino! sources of mRNA. One having ord harv skill in the have been a reasonable expectation of success to co these methods are known in the art and are easy to us

IN RT-PCR products using fluorescent and strand cDNA was synthesized in the : celumn, last paragraph).

we kill in the art at the time the invention or more sources of mRNA using a ected from T3, T7, and SP6 promoter Ita single cDNA library representing two vs together as suggested by Belyavsky et a tenorescence labeled homopolymeric ex molecules as suggested by Luehrsen Jyavsky et al. would have motivated inthesize 1st strand cDNAs from two at: aing a specific sequence tag selected rand cDNAs from two or more A library representing two or more :. • time the invention was made would t ese methods together because all of

MULTINOMIAL TIME SERIES; SCALING; SPECTRAL ENVELOPE 92-1098 001; BLIND PAPTIVE EQUALIZERS; SIGNALS IN FORM CORRELATED RF NOISE; MAXIMUM-LIK, HOOD LOCALIZATION; ROBUST ALGOLITHM; SENSOR ARRAY DATA; DIRECTION FINDING 92-1840 001; SADDLEPOINT APPROXIMATIONS; ASYMPTOTIC PROPERTIES OF A CONDITIONAL MAXIMUM-LIKELIHOOD ESTIMATOR; EXACT DISTRIBUTION; CANONICAL EXPONENTIAL-FAMILIES

RE Referenced Author | Year | VOL | PG | Referenced Work |(RAU)|(RVL)|(RPG)| (RWK)ALOSH M A |1987 |8 | |261 |J TIME SER ANAL ANDERSON T W |1963 |34 |122 |ANN MATH STAT | 1961 | | | STATISTICAL INFERENC | 1981 | | | | TIME SERIES DATA ANA BILLINGSLEY P BRILLINGER D R EATON M L FAHRMEIR L |1970 | | |MULTIPLE TIME SERIES HANNAN E J | 1981 | | 1114 | STRUCTURAL ANAL DISC | 1975 | 5 | 1248 | J MULTIVARIATE ANAL HECKMAN J J IZENMAN A J JOHN S |1963 |25 |363 |SANKHYA A KARLIN S |1991 |86 |27 |J AM STAT ASSOC | 1978 | 7 | 65 | STOCH P APPL LAI C D | 1980 | 5 | 151 | MULTIVARIATE ANAL | 1979 | 7 | 1381 | ANN STAT LEWIS PAW MAGNUS J R MUIRHEAD R J 1 1982 | | ASPECTS MULTIVARIATE |1985 |47 |528 |J ROY STAT SOC B MET RAFTERY A E | 1959 | | | 246 | PROBABILITY STATISTI ROSENBLATT M STOFFER D S |1991 |86 |461 |J AM STAT ASSOC |1987 |8 | 49 | J TIME SER ANAL STOFFER D S TAVARE S | 1989 | | 1117 | MATH METHODS DNA SEQ TYLER D E | 1981 | 9 | 1725 | ANN STAT | 1991 | 133 | 133 | J MOL EVOL

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L18 ANSWER 7 OF 8 BIOSIS COPYRIGHT 2000 BIOSIS DUPLICATE 4
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1984:336303 BIOSIS AN

BA78:72783 DN

WHISENANT E C

DYNAMICS OF DNA OLIGOMERS. TI

TIDOR B; IRIKURA K K; BROOKS B R; KARPLUS M ΑU

DEP. CHEM., HARVARD UNIV., CAMBRIDGE, MASS. 02138. CS

J BIOMOL STRUCT DYN, (1983 (RECD 1984)) 1 (1), 231-252. SO CODEN: JBSDD6. ISSN: 0739-1102.

BA; OLD FS

LA English

in

The techniques of molecular and harmonic dynamics are used to study the internal mobility of 3 double-stranded DNA hexamers. A 60 ps molecular dynamics simulation and a normal mode description of d(CpGpCpGpCpG)2 in the B conformation charcacterize the atomic fluctuations of this structure. A comparison between the 2 approaches validates the harmonic results at room temperature. Detailed examination of the normal modes indicates that only the low-frequency modes are needed

to determine atomic fluctuations. A harmonic analysis is made of d(CpGpCpGpCpG)2 in the Z conformation and of d(TpApTpApTpA)2

the B conformation using only the low-frequency modes. The atomic fluctuations of the 3 alternating pyrimidine-purine helices are compared and the dependence on conformation and sequence are discussed. The insights which theoretical calculations can provide for the interpretation

of experimental results are explored.

Biochemical Methods - Nucleic Acids, Purines and Pyrimidines CC 10052 Biochemical Studies - Nucleic Acids, Purines and Pyrimidines *10062 Biophysics - Molecular Properties and Macromolecules *10506 External Effects - Temperature as a Primary Variable 10614

Application/Control Number: 09/350,609

Art Unit: 1655

The teachings of Sagerstrom et al. have been summurized previously, supra.

Sagerstrom et al. do not disclose cDNA construction after subtracting 1st strand cDNA (tracer) from tissue A with poly(A)+RNA from tissue B (C iver) and tag sequence.

Belyavsky et al. teach a method of identification an 1 cloning differentially expressed mRNA. Figure 1 shows one version of implementing the invention by means of the formation of a set of 3' end labeled fragments of cDNA, dividing it into subjets of fragments with the aid of immobilization on a solid support and sequential treatment—ith a series of restriction nucleases, and separation of the resulting subsets by electrophoresis (column 4). Note that 1st strand cDNA was synthesized by a hemi-tailed (T)₁₃-bio primer and 2nd strand cDNA was synthesized by a homopolymeric dG oligonucleotide tail to the 3' termini of the heteroduplex molecules. In example, synthesis of the second chain of cDNA is done in a reaction mixture containing hybrid mRNA-cDNA, 10 pmol (C)-primer (sequence 5'-AAGGAA i T(C)₁₃), dATP, dGTP, dCTP, dTTP (0.1 mM each) and 1.5 U DNA polymerase Bio-Taq (Biom ster, Russia). The adaptor is added and ligated. Reamplification of the cDNA fragments with the aid of PCR is done using Bio-(T₁₃) primer and a sequence specific primer (column 8). The specific sequence "AAGGAATT" of (C)₁₃ primer can be cut by several different restriction enzy: es.

Belyavsky et al. do not disclose the subtraction of 1: strand cDNA (tracer) from tissue A with poly(A)+ RNA from tissue B (driver) and tag sements selected from T3, T7, and SP6 promoter sequence.

The teachings of Logel et al. have been summarized previously, supra.

Movement 12100

Temperature: Its M. surement, Effects and Regulatio General

Measurement

and Methods 23001

IT Miscellaneous Descriptors
INTERNAL MOBILITY

L18 ANSWER 8 OF 8 BIOSIS COPYRIGHT 2000 BIOSIS DUPLICATE 5

AN 1983:160235 BIOSIS

DN BA75:10235

- TI PERIODICITIES OF DI NUCLEOTIDE SELF INFORMATION VALUES IN PHAGE PHI-X-174 DNA.
- AU FURLONG N B; BECKNER C F
- CS DEP. TUMOR BIOCHEM., UNIV. TEX. SYSTEM CANCER CENT. TUMOR INST., M.D. ANDERSON HOSP., 6723 BERTNER AVE., HOUSTON, TEX., USA.
- SO Z NATURFORSCH SECT C BIOSCI, (1982) 37 (3-4), 321-325. CODEN: ZNCBDA. ISSN: 0341-0382.
- FS BA; OLD
- LA English
- The natural DNA sequence of bacteriophage .vphi.X174, when analyzed as a text of dinucleotides, is shown to display an easily detectable degree of non-randomness by the distribution of values of dinucleotide self-information along the sequence. Self-information corresponding to occurrences of dinucleotides separated by a single nucleotide is somewhat higher than the values which preceed or follow it for every third nucleotide position along the sequence. Autocorrelation coefficients of these values display a strong periodicity and harmonic analysis of the values shows a spike at a value of 3. Self-information autocorrelation periodicity is used as a test of the effect of randomizing portions of the sequence. Any 1 or 2 or the 3 nucleotides in each triplet of the sequence can be chosen at random without losing dinucleotide self-information periodicity except when both the 1st and 3rd nucleotide of all of the triplets in the major .vphi.X174 protein reading frame are randomized. Periodicity is also lost when sequences are generated by randomizing triplets. Autocorrelation and harmonic analysis also indicate other less marked periodic features of dinucleotide self-information values of the native sequence; non-random features are suggested at periods of 12, 20 and 24 nucleotides.
- Biochemical Methods Nucleic Acids, Purines and Pyrimidines 10052
 Biochemical Studies Nucleic Acids, Purines and Pyrimidines *10062
 Replication, Transcription, Translation 10300
 Biophysics Molecular Properties and Macromolecules *10506
 Metabolism Proteins, Peptides and Amino Acids 13012
 Metabolism Nucleic Acids, Purines and Pyrimidines 13014
 Genetics of Bacteria and Viruses *31500
 Virology Bacteriophage *33504
- BC Microviridae 02135
- IT Miscellaneous Descriptors

AUTO CORRELATION PERIODICITY TRIPLET PROTEIN READING FRAME

acidic fibroblast growth factor gene, were synthesized with 5' extensions containing promoter sequences for the T7, T3 and SP6 RNA polymerase promoters. A common antisense primer was used with each of the promoter/aFGF primers to prepare PCR-generated DNA fragments (page 604, abstract). Table 1 showed primers containing T3, T7, and SP6 sequences (page 605). Note that T3, T7, and SP6 polymerase were used in this paper (page 609, left column, last paragraph).

It would have been obvious to one having ordinary skill in the art at the time the invention was made to have synthesized 1st strand cDNAs from two or more sources of mRNA using the primers containing either T3 or T7 sequences as suggested by Logel *et al.* and have produced a single cDNA library representing two or more sources of mRNA by combining 1st strand cDNAs together as suggested by Takahash *et al.*. The prior arts provided by Sagerstrom *et al.* and Takahash *et al.* would have motivated one having ordinary skill in the art to test the possibility to combine 1st strand cDNAs from two or more sources of mRNA together in order to produce a single cDNA library representing two or more sources of mRNA. One having ordinary skill in the art at the time the invention was made would have been a reasonable expectation of success to combine these methods together because all of these methods are known in the art and are easy to use.

8. Claims 6-12, 14, and 15 are rejected under 35 U.S.C. 103(a) as being unpatentable over Sagerstrom *et al.*, (Annu. Rev. Biochem. 66, 751-783, 1907) in view of Belyavsky *et al.*, (US Patent 5,814, 445, filed on July 11, 1995), Logel *et al.*, (Biotechnique 13, 604-610), and Luehrsen *et al.*, (Biotechnique 22, 168-174, January 1997).

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ANSWER 1 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS
      2000:2165 BIOSIS
 AN
 DN
      PREV200000002165
      Cleavage of supercoiled DNA by deoxyribonuclease I in solution
 TI
      and at the electrode surface.
      Fojta, Miroslav (1); Kubicarova, Tatiana; Palecek, Emil
 ΑU
      (1) Institute of Biophysics of the Academy of Sciences of the Czech
 CS
      Republic, Kralovopolska 135, CZ-612 65, Brno Czech Republic
      Electroanalysis, (Oct., 1999) Vol. 11, No. 14, pp. 1005-1012.
 SO
      ISSN: 1040-0397.
 DT
      Article
      English
 LA
 SL
      English
      Cleavage of supercoiled DNA by deoxyribonuclease I (DNase I) in
 AB
      solution and at the surface of the mercury electrode was studied
      by means of AC voltammetry. This technique produces peak 3 which is
      produced only by DNAs containing free ends (such as linear
      double-stranded and single-stranded DNAs and open circular
      DNAs) but not by covalently closed circular (ccc) DNAs.
      Formation of a single interruption of the sugar-phosphate backbone in the
      ccc supercoiled (sc) DNA results in formation of peak 3. Peak 1
      is produced by both ccc DNA molecules as well as by DNAs
      containing free ends; changes in height of this peak occur due to
      DNA cleavage. We show that the kinetics of the cleavage of
      DNA in solution and at the electrode surface
      substantially differ suggesting restricted accessibility of the
      surface-confined DNA for the interaction with the enzyme.
      Cleavage of the immobilized DNA is remarkably influenced by the
      potential of the electrode surface. At positively charged
      surface the enzymati c reaction is inhibited in its initial stage while
     moderately negative charges stimulate the cleavage of the immobilized
     DNA by DNase I.
     Genetics and Cytogenetics - General *03502
CC
     Biochemical Methods - General *10050
     Biochemical Studies - General *10060
     Biophysics - General Brophysical Studies *10502
     Major Concepts
ΙT
        Molecular Genetics (Biochemistry and Molecular Biophysics); Methods
and
        Techniques
     Chemicals & Biochemicals
IT
        deoxyribonuclease I [DNase I]: Sigma, enzyme, kinetics; supercoiled
      DNA: analysis, solution
     Methods & Equipment
IT
        AC voltammetry [alternating current voltammetry]:
        Analysis/Characterization Techniques: CB, analytical method; EG&G PAR
        174A Polarographic Analyzer: equipment; agarose gel electrophoresis:
        gel electrophoresis, separation method; enzymatic cleavage reaction:
        Synthesis/Modification Techniques, chemical method; ethidium bromide
        staining: staining method, staining/visualization; mercury
      electrode: equipment, surface charge
     9003-98-9 (DEOXYRIBONUCLEASE I)
RN
     9003-98-9 (DNASE I)
L22 ANSWER 2 OF 23 SCISEARCH COPYRIGHT 2000 ISI (R)
    1999:77356 SCISEARCH
AN
     The Genuine Article (R) Number: 157GZ
GΑ
     Potential-dependent adsorption/desorption of organic adsorbate at HOPG
TI
     electrode and accompanying delamination of graphite surface
```

L22

7. Claims 1-5 are rejected under 35 U.S.C. 103(a) as being unpatentable over Sagerstrom et al., (Annu. Rev. Biochem. 66, 751-783, 1997) in view of Takahash et al., (Genomics 23, 202-210, 1994) and Logel et al., (Biotechnique 13, 604-610).

Sagerstrom *et al.* review progress of subtractive cloning. Note that they compared the methods of subtractive enrichment and positive selection as shown in Figure 7 (page 772). In Figure 7A, 1st strand cDNA (tracer) from tissue A was used to hybridize with poly(A)+ RNA from tissue B (driver). After twice subtraction to remove hybrids, the remaining fraction can be used to clone insert or synthesize probe (page 772).

Sagerstrom *et al.* do not disclose cDNA construction after subtracting strand cDNA (tracer) from tissue A with poly(A)+ RNA from tissue B (driver) and tag sequences.

Takahash *et al.* teach the construction of an equalized cDNA library from mouse embryos. In this study, RNA from ten different stages of mouse ontogenesis were isolated (page 203, left column, third paragraph) and used for cDNA synthesis. Synthesized ds-cDNAs from ten different stages of mouse ontogenesis were mixed and form "S (straight)-cDNA mixture" (page 203, right column, first paragraph).

Takahash et al. do not disclose the subtraction of strand cDNA (tracer) from tissue A with poly(A)+ RNA from tissue B (driver) and tag sequences

Logel *et al.* teach synthesis of cRNA probes from PCR-generated DNA. In this study, they compared RNA polymerase promoter activities in PCR-generated DNA fragments for use in the in vitro transcription of cRNA probes. Sense oligonucleotide primers, specific for the mouse

Art Unit: 1655

Claim Rejections - 35 USC § 112

- 4. The following is a quotation of the second paragraph of 35 U.S.C. 112:
 - The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.
- 5. Claims 1-5 are rejected under 35 U.S.C. 112, second paragraph, as being incomplete for omitting essential steps, such omission amounting to a gap between the steps. See MPEP § 2172.01. The omitted steps are: combining the products of step a) and b) from claim 1.

Claim Rejections - 35 USC § 103

- 6. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CAR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(f) or (g) prior art under 35 U.S.C. 103(a).

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He Y F (Reprint); Wang Y; Zhu G Y; Wang E ΑU CHINESE ACAD SCI, OF NGCHUN INST APPL CHEM, ELECTROATINT CHEM LAB, CS CHANGCHUN 130022, PL PLES R CHINA (Reprint); CHINESE .. CAD SCI, CHANGCHUN INST APPL CHEM, NATL RES & ANALYT CTR ELECTROCHEM & SPECT, CHANGCHUN 130022, PEOPLES R CHINA CYA PEOPLES R CHINA JOURNAL OF THE ELECTROCHEMICAL SOCIETY, (JAN 1999) Vol. 146, No. 1, pp. SO 250-255. Publisher: ELECTROCHEMICAL SOC INC, 10 SOUTH MAIN STREET, PENNINGTON, NJ 08534. ISSN: 0013-4651. Article; Journal $\mathsf{D}\mathsf{T}$ PHYS; ENGI FS English LA REC Reference Count: 34 In situ electrochemical scanning tunneling microscopy, AB alternating current voltammetry, and electrochemical quartz crystal microbalance have been employed to follow the potential-dependent adsorption/desorption processes of nucleic acid bases on highly oriented pyrolytic graphite (HOPG) electrode. The results show that (i) potential-dependent adsorption/desorption of nucleic acid bases on HOPG electrode was accompanied by delamination of the HOPG surface, and the delamination initiates from steps or kinks on the electrode surface, which provide highly active sites for adsorption; (ii) the delamination usually occurred when the electrode potential was changed or when the electrode was at potentials where the phase transition of adsorbate occurred. These results suggest that the surface stress resulting from the interaction between the substrate and adsorbate, as well as the interaction due to potential-induced surface charge distribution and the hysteresis of charge equilibrium are the main . factors resulting in HOPG delamination. (C) 1999 The Electrochemical Society. s0013-4651(97)12-013-4. All rights reserved. ELECTROCHEMISTRY; MATERIALS SCIENCE, COATINGS & FILMS CC STP KeyWords Plus (R): SCANNING-TUNNELING-MICROSCOPY; DIFFERENTIAL CAPACITANCE; PYROLYTIC-GRAPHITE; MONOLAYER GUANINE; AQUEOUS-SOLUTIONS; NACL SOLUTION; STRESS; STM; RECONSTRUCTION; INTERFACE RE

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MULLER J E	1986 56 1583	PHYS REV LETT

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                                          J ELECTROANAL EM
                         272 | 36
                                   |257
RANDIN J P
                         71 | 118 | 711 | J ELECTROCHEM C
RANDIN J P
                       |1992 |272 |318 |SURF SCI
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                       |1959 |155 |206 |Z PHYS
SAUERBREY G
                       |1991 |312 |293
                                          J ELECTROANAL CH INF
SRINIVASAN R
                                   |1464 |J PHYS CHEM-US
                       |1994 |98
TAO N J
                                   |7422 |J PHYS CHEM-US
TAO N J
                       |1994 |98
                                 |4445 |LANGMUIR
TAO N J
                       |1995 |11
                       |1994 |301 |L217
TAO N J
                                          |SURF SCI
                       |1996 |419 |1
WANG Y
                                          | J ELECTROANAL CHEM
                       |1996 |364 |L530 |SURF SCI
ZHANG J D
L22
    ANSWER 3 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS
                                                        DUPLICATE 1
AN
     1999:247083 BIOSIS
DN
     PREV199900247083
TI
     DNA-modified electrodes Part 3.: Spectroscopic
     characterization of DNA-modified gold electrodes.
     Zhao, Yuan-Di; Pang, Dai-Wen (1); Hu, Shen; Wang, Zong-Li; Cheng, Jie-Ke;
AU
     Qi, Yi-Peng; Dai, Hong-Ping; Mao, Bing-Wei; Tian, Zhong-Qun; Luo, Jin;
     Lin, Zhong-Hua
     (1) Department of Chemistry, Wuhan University, Wuhan, 430072 China
CS
     Analytica Chimica Acta, (May 3, 1999) Vol. 388, No. 1-2, pp. 93-101.
SO
     ISSN: 0003-2670.
    Article
DT
    English
LA
\mathtt{SL}
     English
     DNA-modified gold electrodes were characterized by
AΒ
     scanning tunneling microscopy (STM), Raman spectroscopy, in situ UV/Vis
     reflection spectroscopy, X-ray photoelectron spectroscopy (XPS) and
     alternating current (AC) impedance. It has been found
     that dsDNA adsorbed firmly on gold surfaces lies strand-on in an ordered
     saturated monolayer, and ssDNA strands exist in a honeycomb-like form on
     the surfaces. The bases and phosphate groups of DNA backbone
     interacting with gold electrode surfaces play an important role
     in DNA immobilization onto gold electrode surfaces.
CC
     Biochemical Methods - General *10050
     Comparative Biochemistry, General *10010
     Biochemical Studies - General *10060
     Biochemical Studies - Nucleic Acids, Purines and Pyrimidines *10062
     Biochemical Studies - Minerals *10069
     Biophysics - General Biophysical Studies *10502
     Biophysics - Molecular Properties and Macromolecules *10506
     Biophysics - Bioengineering *10511
    Major Concepts
        Biochemistry and Molecular Biophysics; Equipment, Apparatus, Devices
        and Instrumentation; Methods and Techniques
     Chemicals & Biochemicals
{	t IT}
        gold; DNA: Sino-American Biotechnical, immobilization,
        purification
    Methods & Equipment
IT
        scanning tunneling microscopy: microscopy method, tunneling
microscopy;
       AC impedance measurement: Detection/Labeling Techniques, analytical
       method; DNA-modified gold electrodes: applications,
        characterization, laboratory equipment; LabRam I Confocal MicroRaman
        system: Dilor, equipment; Raman spectroscopy: analytical method,
        spectroscopic techniques: CB; UV/Vis reflection spectrophotometer:
       equipment; UV/Vis reflection spectroscopy: analytical method,
        spectroscopic techniques: CB; VG ESCA-LAB MKII spectrometer:
laboratory
       equipment; X-ray photoelectron spectroscopy: analytical method,
        spectroscopic techniques: CB
    Miscellaneous Descriptors
ΙT
       electrochemistry
```

Wu, Jin-Tian,; Huang, Yin; Zhou, Jian-Zhang; Luo, Jin; Lin, Zhong-Hua (1)

Bioelectrochemistry and Bioenergetics, (Nov., 1997) Vol. 44, No. 1, pp.

(1) State Key Lab. Physical Chem. Solid Surfaces, Dep. Chem., Inst.

Physical Chem., Xiamen Univ., Xiamen 361005 China

TI

AU

CS

SO

electrode.

151-154.

ISSN: 0302-4598.

```
Article
\mathtt{DT}
     English
LA
    DNA was studied by ans of cyclic voltammetry (CV)
AB
     combination with a mercury film electrode (MFE) using
     conventional CV, differential pulse voltammetry (DPV), alternating
     current voltammetry (ACV). The MFE is sufficiently stable and can
     be used to study electrochemical behaviors of DNA in negative
     potential region. This means that MFE is ready to be one kind of solid
     electrode at which more useful electrochemical techniques can be
     carried out, such as spectroelectrochemical techniques, Redox characters
     of DNA treated by pure perchloric acid (HClO4) was studied at
     MFE. It seems that pure HClO4 would not only bring about the denaturation
     of DNA but the degradation of it. Pure HClO4 is not suitable for
     performing the denaturation of DNA.
     Biochemical Methods - Nucleic Acids, Purines and Pyrimidines *10052
CC
     Biochemical Studies - Nucleic Acids, Purines and Pyrimidines
                                                                    *10062
     Biophysics - General Biophysical Techniques *10504
     Major Concepts
IT
        Methods and Techniques; Molecular Genetics (Biochemistry and Molecular
        Biophysics)
     Chemicals & Biochemicals
IT
        DNA: electrochemical behavior
     Methods & Equipment
IT
        alternating current voltammetry: analytical method;
        cyclic voltammetry: analytical method; differential pulse voltammetry:
        analytical method; mercury film electrode: equipment
     7439-97-6 (MERCURY)
RN
L22 ANSWER 6 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS
     1995:224095 BIOSIS
AN
     PREV199598238395
DN
     Voltammetry of adsorbed cancerostatic actinomycins.
TI
     Ibrahim, M. S. (1); Ahmed, Z. A.; Temerk, Y. M.; Berg, H.
AU
     (1) Chem. Dep., Fac. Sci., Assiut Univ., Assiut Egypt
CS
     Bioelectrochemistry and Bioenergetics, (1995) Vol. 36, No. 2, pp.
SO
149-156.
     ISSN: 0302-4598.
     Article
DT
     English
\mathtt{L}\mathtt{A}
     A systematic study of the adsorption and association of the cancerostatic
AB
     drug actinomycin-C-1 (ACT) at a hanging mercury drop electrode
     (HMDE) has been conducted using phase-sensitive a.c. voltammetry and
     cyclic voltammetry (CV). At all bulk concentrations, the adsorbed layer
is
     transformed into a condensed film by the significant stacking forces
     acting between adjacent rings of the phenoxazone residues. The nucleation
     and growth mechanism is confirmed and the data are analysed using the
     Avrami equation. The adsorption parameters for the condensed film were
     evaluated at various pH values. In addition, the preparative
     electrochemical reduction of ACT was performed using the large-scale
     electrolysis and differential pulse polarography. The consequences for
     DNA interaction and membrane adsorption are discussed.
     Biochemical Studies - General *10060
CC
     Biochemical Studies - Nucleic Acids, Purines and Pyrimidines
                                                                     10062
     Biochemical Studies - Minerals
                                     10069
     Biophysics - Molecular Properties and Macromolecules *10506
     Biophysics - Membrane Phenomena *10508
     Pharmacology - General *22002
     Neoplasms and Neoplastic Agents - Therapeutic Agents; Therapy *24008
     Major Concepts
IT
        Biochemistry and Molecular Biophysics; Membranes (Cell Biology);
        Pharmacology; Tumor Biology
     Chemicals & Biochemicals
IT
        ACTINOMYCINS; ACTINOMYCIN C1; PHENOXAZONE; MERCURY
     Miscellaneous Descriptors
IT
```

A systematic study on the adsorption and association of 6-thiopurine AB(6-TP) and 6-thiopurine riboside (6-TPR) has been carried out at various pH values and the adsorption parameters were determined quantitatively. The adsorption was followed by out-of-phase alternating current voltammetry and cyclic voltammetry at a hanging mercury drop electrode. A comparative study was undertaken on the adsorption and association of the investigated thiopurines and the similar

type of nucleic acid components containing purine bases. The base-containing thio group enhances stacking interaction and facilitates formation of the perpendicularly stacked layer on the electrode surface.

Medical Descriptors: CT

*adsorption

article

ph

potentiometry

priority journal

Drug Descriptors:

*mercaptopurine: AN, drug analysis

6 thiopurine riboside: AN, drug analysis

unclassified drug

(mercaptopurine) 31441-78-8, 50-44-2, 6112-76-1 RN

Sigma (United States) CO

ANSWER 8 OF 23 SCISEARCH COPYRIGHT 2000 ISI (R) L22

93:74543 SCISEARCH NA

The Genuine Article (R) Number: KK281 GΑ

ELECTROPORATION OF INOSITOL 1,4,5-TRIPHOSPHATE INDUCES REPETITIVE CALCIUM TIOSCILLATIONS IN MURINE OOCYTES

RICKORDS L F; WHITE K L (Reprint) ΑU

UTAH STATE UNIV, CTR BIOTECHNOL, DEPT ANIM DAIRY & VET SCI, LOGAN, UT, CS 84322

CYA USA

JOURNAL OF EXPERIMENTAL ZOOLOGY, (01 FEB 1993) Vol. 265, No. 2, pp. SO 178-184.

ISSN: 0022-104X.

Article; Journal \mathtt{DT}

FS LIFE; AGRI

ENGLISH LA

REC Reference Count: 39

The purpose of se experiments was to determine the effect of electroporation of 3 into the cytosol of murine see ndary occytes and AB evaluate any alterations in [Ca2+]i resulting from Ca2+ release from intracellular stores. In addition, we evaluated the effect of ethanol (ETOH) on the release of Ca2+ from intracellular stores. Oocytes were loaded with the Ca2+ indicator fluo-3 by incubation in 100 mul drops of medium containing 2 muM fluo-3/AM for 60 min at 37-degrees-C. Changes in fluorescence were monitored by use of an inverted microscope which had been connected to a spectrofluorometer. Fluorescent intensity measurements

were acquired for a minimum of 416 sec time span or up to 1,248 sec, with integration readings of 1 sec duration obtained every 2 sec throughout the

measurement period. The experimental design consisted of comparing the rise in [Ca2+]i of fluo-3 loaded secondary oocytes subjected to electroporation in PBS and Ca2+-free PBS, each containing 25 muM IP3, to that elicited by PBS and Ca2+-free PBS containing a final concentration

7% ETOH. Non-pulsed control secondary oocytes were placed in PBS + 25 muM IP3 during monitoring of [Ca2+]i fluorescence. Pulsed control secondary oocytes were placed in Ca2+-free PBS, subjected to electroporation pulse, and monitored for [Ca2+]i fluorescence.

Electroporation of IP3 was accomplished by placing fluo-3 loaded secondary oocytes between the electrodes of a glass slide fusion chamber which had been overlaid with 300 mul of PBS + 25 muM IP3 or Ca2+-free PBS + 25 muM IP3. A 5 sec, 3 volt, alternating current (AC) alignment pulse followed by a single, square wave, direct current (DC) fusion pulse of 1.56 kV.cm-1 for 99 musec was applied to the electrodes. For ETOH treatment, fluo-3 loaded secondary oocytes were placed in PBS or Ca2+-free PBS and allowed to equilibrate

7 min in the dark. No pulse was applied to ETOH treatment secondary oocytes. Micropipets were used to keep the secondary oocyte in a fixed position throughout the measurement period. After a 20 sec baseline fluorescent reading was obtained, fluorescent measurement was interrupted and 150 mul of PBS (or Ca2+-free PBS) was removed and replaced with 150 mul of 14% ETOH in PBS (or Ca2+-free), bringing the final concentration after equilibration to 7% ETOH. Fluorescent intensity measurement resumed immediately following the addition of 14% ETOH. A dramatic and immediate rise in [Ca2+]i was observed upon application of electropolation pulse

[Ca2+]i was maintained at an elevated level for a minimum of 14 min. Repetitive [Ca2+]i oscillations were obtained in mouse secondary oocytes after electroporation of 25 muM IP3 in Ca2+-free PBS that occurred for 20.5 min with a gradual increase in the interval between [Ca2+]i oscillation peaks over time. After ETOH treatment, a dramatic rise in mouse secondary oocyte [Ca2+]i in PBS and Ca2+-free PBS was observed. There was no significant different (P > 0.05) in [Ca2+]i between PBS + ETOH and Ca2+-free PBS + ETOH, indicating the rise in [Ca2+]i resulted from a release of Ca2+ from intracellular stores. The ability to consistently produce repetitive [Ca2+]i oscillations may aid in the study of post-fertilization development and cell cycle events. Current studies are being conducted to determine if IP3 can be used to enhance the rate

electric pulse induced parthogenesis and subsequent development. ZOOLOGY CC

STP KeyWords Plus (R): GOLDEN-HAMSTER EGGS; SEA-URCHIN EGGS; INTRACELLULAR FREE CALCIUM; HYPERPOLARIZING RESPONSES; PERIODIC INCREASE; BINDING PROTEIN; ELECTRIC-FIELDS; XENOPUS-LAEVIS; DNA-SYNTHESIS; FERTILIZATION

92-6934 002; TRANSMITTER RELEASE INCREASES INTRACELLULAR CALCIUM; RFINOSITOL

TRISPHOSPHATE IN XENOPUS OOCYTES; MOUSE THYMOCYTES 92-2219 001; PROTEIN-KINASE-C ISOFORMS; PHORBOL ESTER; CULTURED RAT

for

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of

MESANGIAL CELLS				
RE Referenced Author (RAU)	ar (RPY)	VOL (RVL)	PG (RPG)	Referenced Work (RWK)
	+=====	+====·	+=====: . 500	+=====================================
BERRIDGE M J	'	403	589 315	NATURE
BERRIDGE M J	1984	•	315 677	J CELL BIOL
BUSA W B	•	•	•	GAMETE RES
COLONNA R	11989	124	171	NATURE
CUTHBERTSON K S R	11981	•	1754	NATURE
CUTHBERTSON K S R	1985	316	541	CALCIUM CELL FUNCTIO
EPEL D	11982	12	355 1	CELL DIFFER DEV
EPEL D	11990	129	•	CURR TOP DEV BIOL
EPEL D	1978	12	1185	·
HAN J K	1990	110	1103	
IGUSA Y	11983	1340	611	J PHYSIOL-LONDON
IGUSA Y	1983	340	633	J PHYSIOL-LONDON
IGUSA Y	1986	377	193	J PHYSIOL-LONDON
JAFFE L A	1983	196	317	DEV BIOL
JAFFE L F	1985	}	1127	BIOL FERTILIZATION
JAFFE L F	1983	199	1265	DEV BIOL
KAO J P Y	1989	264	8179	J BIOL CHEM
KAUFMAN M H	1983		120	EARLY MAMMALIAN DEV
KAUFMAN M H	1982	171	139	J EMBRYOL EXP MORPH
MCCULLOH D H	1983	•	1372	DEV BIOL
MINTA A	1989	264	18171	J BIOL CHEM
MIYAZAKI S	1991	12	205	CELL CALCIUM
MIYAZAKI S	1986	118	1259	DEV BIOL
MIYAZAKI S	1988	106	345	J CELL BIOL
MIYAZAKI S	11988	50	390	J PHYSL SOC JAPAN
MIYAZAKI S	11981	1290	1702	NATURE
MIYAZAKI S	1982	179	931	P NATL ACAD SCI USA
OZAWA H	11989	138	477	J CELL PHYSIOL
OZIL J P	11990	1109	117	DEVELOPMENT
RICKORDS L F	11992	31	152	MOL REPROD DEV
RODAN G A	11978	1199	1690	SCIENCE
STEINHARDT R A	1988	•	364	NATURE
SWANN K	11990	•	1295	DEVELOPMENT
SWANN K	11986	•	2333	J CELL BIOL
TURNER P R	11986	•	170	J CELL BIOL
WHITAKER M	•	•	525	•
11	•	•	636	·
WHITAKER M J	11989		1459	
WOOD M J	11987	•	1255	MAMMALIAN DEV PRACTI
WOOD II O	12007	1	,	
1.22 ANSWER 9 OF 23 BI	OSIS	COPYRI	GHT 200	00 BIOSIS DUPLICAT

- L22 ANSWER 9 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS DUPLICATE 5
- 1990:517072 BIOSIS AN
- BA90:134348 DN
- ALTERNATING CURRENT VOLTAMMETRIC DETERMINATION OF TIDNA DAMAGE.
- KRZNARIC D; COSOVIC B; STUEBER J; ZAHN R K ΑU
- CENT. MARINE RES. ZAGREB, RUDER BOSKOVIC INST., BIJENICKA 54, 41000 CS ZAGREB, YUGOSL.
- CHEM-BIOL INTERACT, (1990) 76 (1), 111-128. SO CODEN: CBINA8. ISSN: 0009-2797.
- BA; OLD FS
- English LA
- The conditions for alternating current (a.c.) ABvoltammetric DNA determinations have been investigated with respect to its use with alkaline filter elution techniques at low DNA concentrations. In inorganic electrolyte solutions three current peaks can be distinguished: peak I around -1.1 V caused by the reorientation or desorption of DNA segments; peak II around -1.2 V caused by the native DNA (nDNA) form; peak III caused by denatured DNA (dDNA) at -1.4 V. Sonication of nDNA increases the

peak current, however not with dDNA. Both dDNA and nDNA give linear peak current increments the DNA increments, their regression lines cutting the concent tion axis at the origin. In filter elution techniques

organic bases are often used. Adding ethanolamine (EA) elution buffer decreases the peak amplitude of DNA. It turns out that an unknown substance, perhaps a protein or RNA, elutes from the filters and gives rise to a current peak at about -1.3 V. This substance can interfere with the dDNA by competing for electrode surface area, since it diffuses much faster than the large molecules of the DNA. Since however, dDNA has a higher affinity for the electrode surface, after enough time, usually few minutes, the dDNA increasingly displaces the substance and occupies the surface. The same is valid for other organic molecules and thus also for EA. It is therefore remarkable that the unknown substance can be altered by ultrasonication, so that it will no longer interfere with dDNA, in contrast to EA. EA, on the other hand, can be "titrated". When EA is present at short accumulation times it prevents dDNA adsorption. By

adding

dDNA, the EA can be scavenged and further addition will adsorb and thus increase peak current in proportion to the concentration of the DNA present. The conditions for voltammetric DNA determination have been investigated obeying the recognized interactions. Avoiding organic bases and using inorganic ones would simplify the determination procedure. The reproducibility of the procedure in the range

of 50-60 ng DNA/ml has been found to be .+-. 6%.

Genetics and Cytogenetics - Animal *03506

Biochemical Methods - Nucleic Acids, Purines and Pyrimidines 10052

Biochemical Studies - Nucleic Acids, Purines and Pyrimidines *10062

Biophysics - Molecular Properties and Macromolecules *10506

External Effects - Electric, Magnetic and Gravitational Phenomena *10610

External Effects - Physical and Mechanical Effects *10612

IT Miscellaneous Descriptors HOLOTHURIA-TUBULOSA

L22 ANSWER 10 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS

AN 1990:127494 BIOSIS

DN BA89:66305

TI CYCLIC VOLTAMMETRY OF METAL-POLYELECTROLYTE COMPLEXES COMPLEXES OF CADMIUM

AND LEAD WITH DNA.

AU SEQUARIS J-M; ESTEBAN M

CS INST. APPLIED PHYSICAL CHEM., NUCLEAR RES. CENTER KFA JUELICH, P.O. BOX 1913, D-5170 JUELICH, WEST GERMANY.

SO ELECTROANALYSIS, (1990) 2 (1), 35-42. CODEN: ELANEU. ISSN: 1040-0397.

FS BA; OLD

LA English

of

Cyclic voltammetry was used for the determination of the association constants of Pb2+ and Cd2+ with deoxyribonucleic acid. The adsorption of the biological polyelectrolyte at the mercury electrode surface was controlled by the alternating current voltammetric method, which permits corrective factors to be introduced in the evaluation of cyclic voltammetric responses. The results are based on the analysis of the labile complexation of the slow-diffusing DNA by studying the current intensity peak as well as the peak potential shift

Pb2+ and Cd2+. The association constants (.beta.) obtained from the two treatments are in satisfactory agreement. The dependence of the conditional association constant (.beta.1) for the Cd-DNA system on the monovalent ion (Na+) concentration is also reported.

CC Biochemical Methods - General *10050

Biochemical Methods - Nucleic Acids, Purines and Pyrimidines *10052

Biochemical Methods - Minerals *10059

Biochemical Studies - Nucleic Acids, Purines and Pyrimidines 10062 Biochemical Studie Minerals 10069 Biophysics - Genera Biophysical Techniques *10504 Biophysics - Molecular Properties and Macromolecules *10506 External Effects - Electric, Magnetic and Gravitational Phenomena *10610 Miscellaneous Descriptors IT CONDITIONAL ASSOCIATION CONSTANT ASSOCIATION CONSTANT CURRENT INTENSITY PEAK PEAK POTENTIAL SHIFT 7439-92-1 (LEAD) RN 7440-43-9D (CADMIUM) L22 ANSWER 11 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS 1988:151829 BIOSIS AN DN BA85:75482 ELECTRIC FIELD EFFECTS IN NUCLEIC ACIDS ADSORPTION OF TIADENINE AT THE NEGATIVELY CHARGED ELECTRODE. VETTERL V; JANCAR J; ZALUDOVA R ΑU INST. BIOPHYS., CZECH. ACAD. SCI., 612 65 BRNO, CZECH. CS FOLIA FAC SCI NAT UNIV PURKYNIANAE BRUN BIOL, (1987) 0 (85), 85-94. SO CODEN: FFUBAP. BA; OLD FS English LA The alternating current polarograms of adenine at ABdifferent pH were measured. With increasing concentration of adenine a sort of pit appears on the a.c. polarograms near the potential of the electrocapillary maximum, indicating the region of potentials at which the adsorbed adenine molecules associate. Besides the pit occurring in the vicinity of the electrocapillary maximum potential a more negative pit around the potential of -1.2 V was observed in the pH range 3.9-5.5 at high concentrations of adenine. This more negative pit corresponds to the association of adenine molecules electrostatically adsorbed to the mercury surface. Biochemical Studies - Nucleic Acids, Purines and Pyrimidines *10062 CC Biophysics - General Biophysical Techniques 10504 Biophysics - Molecular Properties and Macromolecules *10506 External Effects - Electric, Magnetic and Gravitational Phenomena *10610 Miscellaneous Descriptors ITALTERNATING CURRENT POLAROGRAM 73-24-5 (ADENINE) RN ANSWER 12 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS DUPLICATE 6 1987:61960 BIOSIS NABA83:30286 DNALTERNATING CURRENT VOLTAMMETRIC DETERMINATION OF \mathtt{TI} DNA CONCENTRATIONS AT A MICROGRAM PER LITER LEVEL. KRZNARIC D; COSOVIC B ΑU CENT. MARINE RES. ZAGREB, RUDJER BOSKOVIC INST., ZAGREB, YUGOSLAVIA. CS ANAL BIOCHEM, (1986) 156 (2), 454-462. SO CODEN: ANBCA2. ISSN: 0003-2697. BA; OLD FS English LAAlternating current voltammetry is used as a fast and ΆB highly sensitive method of DNA detection, at a microgram per liter level. The method is based on the measurement of adsorption effects of denatured DNA at the hanging mercury drop electrode . The proposed procedure consists of thermal denaturation of DNA , which is followed by electrochemical detection of denatured DNA . A sharp adsorption peak of denatured DNA, at the potential of -1.4 V, is measured in 0.3 mol/liter NaCl and 0.03 mol/liter NaHCO3 (pH about 9) after an accumulation of DNA at the electrode surface. To enhance the sensitivity, the solution is stirred during

adsorption. The influence of proteins, a polysaccharide, and RNA on the DNA determination was also studied. Biochemical Methods Nucleic Acids, Purines and Pyr Aidines *10052 CC Biochemical Studies - Nucleic Acids, Purines and Pyrimidines 10062 Biochemical Studies - Proteins, Peptides and Amino Acids 10064 Biochemical Studies - Carbohydrates 10068 Biophysics - General Biophysical Techniques *10504 L22 ANSWER 13 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS 1982:233058 BIOSIS ANBA74:5538 DN POLAROGRAPHY OF CIRCULAR DNA. TIVOJTISKOVA M; LUKASOVA E; JELEN F; PALECEK E ΑU INST. BIOPHYSICS, CZECHOSLOVAK ACADEMY OF SCI., KRALOVOPOLSKA 135, 612 CS 65, BRNO, CZECHOSLOVAKIA. BIOELECTROCHEM BIOENERG, (1981) 8 (5), 487-496. SO CODEN: BEBEBP. ISSN: 0302-4598. BA; OLD FS English LAClosed duplex (cd) and open circular (oc) forms of DNA of the AB plasmid Col El were studied by means of AC and differential pulse polarography (dpp). Adsorption properties of oc DNA (at pH 8) agreed in principle with those of linear DNA, cd DNA was less firmly adsorbed at the dme (dropping mercury electrode), compared with oc DNA. At low ionic strengths cd DNA was adsorbed at potentials more positive than the pzc via unscreened, negatively charged phosphates, and around -0.55 V (vs. sce (saturated calomel electrode)) it produced a much higher tensammetric peak than oc DNA. At moderate ionic strengths oc DNA produced a well-developed peak 1 at about -1.1 V. Peak I of cd DNA was considerably smaller, in accord with a much weaker adsorption of this DNA at a potential more negative than the pzc. Under conditions suitable for the polarographic reduction of single-stranded DNA, cd DNA behaved as non-reducible, as detected by the absence of dpp peaks in the potential region from -1.3 to -1.5 V. oc DNA produced dpp peak II, so far observed only with linear double-stranded DNA. Thermally denatured oc DNA produced a high peak III characteristic for denatured DNA. A dpp method for the determination of oc DNA in samples of cd DNA was designed. The experimental data obtained were utilized for explaining the role of bases in the interaction of a polynucleotide molecule with the dme and for elucidating some changes in DNA conformation in the bulk Biochemical Methods - Nucleic Acids, Purines and Pyrimidines *10052 CC Biochemical Studies - Nucleic Acids, Purines and Pyrimidines *10062 Biophysics - General Biophysical Techniques *10504 Biophysics - Molecular Properties and Macromolecules 10506 Physiology and Biochemistry of Bacteria *31000 Genetics of Bacteria and Viruses 31500 Microbiological Apparatus, Methods and Media 32000 Enterobacteriaceae 04810 BC Miscellaneous Descriptors ITCOL-E-1 PLASMID CLOSED DUPLEX DNA OPEN CIRCULAR DNA SINGLE STRANDED DNA DOUBLE STRANDED DNA DENATURED DNA LINEAR DNA INTERCALATION CONFORMATION ALTERNATING CURRENT POLAROGRAPHY DIFFERENTIAL PULSE POLAROGRAPHY L22 ANSWER 14 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS 1980:200942 BIOSIS ANBA69:75938 DNINTERACTION OF NUCLEIC-ACIDS WITH ELECTRICALLY CHARGED

SURFACES 7. THE EFFECT OF IONIC STRENGTH OF NEUTRAL MEDIUM ON THE

TI

CONFORMATION OF DNA ADSORBED ON THE MERCURY ELECTRODE. BRABEC V ΑU INST. BIOPHYS., CZE. ACAD. SCI., 61265 BRNO, CZECA CS BIOPHYS CHEM, (1980) 11 (1), 1-8. SO CODEN: BICIAZ. ISSN: 0301-4622. BA; OLD FS English LATriangular-wave direct current (DC) voltammetry at a hanging mercury drop AB electrode and phase-selective alternating current (AC) polarography at a dropping mercury electrode were used for the investigation of adsorption of double-helical (ds) DNA at mercury electrode surfaces from neutral solutions of 0.05-0.4 M HCOONH4. It was found for the potential region T (from -0.1 V up to approximately -1.0 V) that the height of voltammetric peaks of ds DNA is markedly influenced by the initial potential only at

A decrease of differential capacity (measured by means of AC polarography) in the region T depended markedly on the electrode potential only at relatively low ionic strength. The following conclusions were made

concerning the interaction of ds DNA with a mercury electrode charged to potentials of the region T in neutral medium of relatively low ionic strength (.mu. < 0.3). When ds DNA is adsorbed, a significantly higher number of DNA segments is anchored in the positively charged electrode surface than in the surface bearing a negative charge. In the region T, especially adsorbed labile regions of ds DNA are opened in the electrode surface, which are present in ds DNA already in the bulk of the solution. In the narrow region of potentials in the vicinity of the zero charge potential a higher number of ds DNA segments can be opened, probably as a consequence of the strain which could act on the ds DNA molecule in the course of the segmental adsorption/desorption process.

relatively low ionic strength (.mu.) (from 0.05 up to approximately 0.3).

Biochemical Methods - Nucleic Acids, Purines and Pyrimidines 10052 CC Biochemical Studies - Nucleic Acids, Purines and Pyrimidines *10062 Biophysics - General Biophysical Techniques 10504 Biophysics - Molecular Properties and Macromolecules *10506

7439-97-6 (MERCURY) RN

DUPLICATE 7 L22 ANSWER 15 OF 23 BIOSIS COPYRIGHT 2000 BIOSIS

1978:187819 BIOSIS NA

BA66:316 DN

- ALTERNATING CURRENT POLAROGRAPHIC INVESTIGATION OF TIPOLY SACCHARIDES IN DNA.
- MALFOY B; REYNAUD J A ΑU
- CENT. BIOPHYS. MOL., 45045 ORLEANS CEDEX, FR. CS
- ANAL BIOCHEM, (1978) 84 (1), 1-11. SO CODEN: ANBCA2. ISSN: 0003-2697.
- BA; OLD FS
- English $_{
 m LA}$
- Polysaccharides alone or in the presence of DNA are studied by ABAC polarography. When neutral and basic polysaccharides are used, the polarograms recording the quadratic component of the current display 1 capacitive peak at -1650 mV (SCE [saturated calomel electrode]). Acid polysaccharides never show this peak and are desorbed from the electrode at more positive potentials. If dextran is used as a reference, this peak allows the determination of the amount of neutral polysaccharides in solution up to 2 .mu.g/ml. The height of this peak has no relation to the ionic strength or pH of the solution within the investigated range. The concentration and MW of DNA enclosed in the solution exert no influence on the peak height. The presence of polysaccharides causes DNA peaks to decrease considerably. AC polarography can be regarded as a quick, convenient and sensitive method for performing the titration of polysaccharides alone or mixed with

```
DNA.
    Biochemical Method Nucleic Acids, Purines and Pyridines Biochemical Method Carbohydrates *10058
                                                                    *10052
CC
     Biochemical Studies - Nucleic Acids, Purines and Pyrimidines 10062
     Biochemical Studies - Carbohydrates 10068
     Biophysics - General Biophysical Techniques *10504
L22 ANSWER 16 OF 23 CAPLUS COPYRIGHT 2000 ACS
     1975:125551 CAPLUS
NA
     82:125551
DN
     Adsorption of DNA at the mercury-electrolyte interface. V.
ΤI
     Influence of temperature on the structure of the adsorption layer of
     DNA
     Flemming, J.
ΑU
     Zentralinst. Mikrobiol. Exp. Ther., DAW, Jena, E. Ger.
CS
     Stud. Biophys. (1974), 45,, 21-7
SO
     CODEN: STBIBN
     Journal
DT
LA English
    33-7 (Carbohydrates)
CC
     Section cross-reference(s): 22
     The temp. influence on the adsorption layer of DNA at the
AB
     interface between a hanging Hg drop electrode and a buffered aq.
     NaCl soln. was detd. via alternating current
     polarography and the differential capacity was measured at several
     potentials. The extending of adsorbed DNA mols. into soln.
     increased with increasing temp. and from these measurements the
premelting
     and denaturation of DNA can be estd.
     DNA adsorption temp dependence; mercury electrode
\mathtt{ST}
     DNA adsorption; polarography DNA adsorption
     Deoxyribonucleic acids
ΙT
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
        (adsorption of, effect of temp. on)
     Adsorption
{	t IT}
         (of deoxyribonucleic acids, effect of temp. on)
L22 ANSWER 17 OF 23 EMBASE COPYRIGHT 2000 ELSEVIER SCI. B.V.
AN
     75001532 EMBASE
    1975001532
DN
     The relation between adsorbability and polarographic reducibility of
TI
     single stranded polynucleotides.
     Brabec V.; Palecek E.
ΑU
     Inst. Biophys., Czech. Acad. Sci., Brno, Czechoslovakia
ÇS
     Studia Biophysica, (1974) 42/1 (1-6).
     CODEN: STBIBN
     Journal
DT
             Clinical Biochemistry
     029
FS
     English
LA
     Interactions of single stranded polycytidylic acid (poly(C)) with mercury
AΒ
     electrode were followed by means of direct current (dc) and
      alternating current (ac) polarography. It was found that
      the polarographic reduction of poly (C) takes place only in the adsorbed
      state. The reduction limiting currents of poly (C) exhibited properties
      typical for adsorption currents in agreement with the above finding.
      Different shapes of dc polarographic curves of poly (C) could be
explained
      in the same way as those of denatured DNA and polyadenylic acid
      (poly (A)) by inhibition of reduction current due to polynucleotide
      desorption from the negatively charged surface of mercury
      electrode. The character of the electrode process which
      is responsible for reduction of poly (C) on mercury electrode
      was similar to the character of the process at which denatured DNA
      and poly (A) were reduced.
      Medical Descriptors:
 CT
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in vitro study theoretical study methodology Drug Descriptors: *dna *polycytidylic acid *polynucleotide (dna) 9007-49-2; (polycytidylic acid) 30811-80-4 RN L22 ANSWER 18 OF 23 CAPLUS COPYRIGHT 2000 ACS 1972:430499 CAPLUS AN 77:30499 DN Interactions of polynucleotides with the mercury electrode TIBrabec, V.; Palecek, E. AU Inst. Biophys., Czech. Acad. Sci., Brno, Czech. CS Proc. Conf. Appl. Phys. Chem., 2nd (1971), Volume 1, 523-7. Editor(s): SO Buzas, Ilona. Publisher: Akad. Kiado, Budapest, Hung. CODEN: 24IUAO Conference DTEnglish LA6-2 (General Biochemistry) CC Section cross-reference(s): 9 Adsorption of polynucleotides was studied by means of Breyer AB alternating current (a.c.) polarog. According to the a.c. polarog. behavior of various model compds. a scheme of adsorption of polynucleotides was suggested. In medium of higher ionic strength, when the charges of phosphate groups of DNA were screened by ions of the electrolyte, double-helical DNA was adsorbed as an electroneutral substance. Under low ionic strength, the segment of double-helical DNA in which all the charges of phosphate groups were not screened were adsorbed on the pos. electrode surface. Single-stranded polynucleotides were adsorbed on the Hg electrode mainly through bases. mercury electrode polynucleotide interaction STChains, chemical IT(helical conformation of, of polynucleotides, mercury electrode interaction in relation to) Electrodes IT(mercury, adsorption of polynucleotides, helical conformation in relation to) Adsorption IT (of polynucleotides on mercury electrode, helical conformation in relation to) Nucleotides, properties IT RL: PRP (Properties) (poly-, adsorption on mercury electrode, helical conformation in relation to) Ions in liquids ${
m IT}$ (strength of, polynucleotide adsorption on mercury electrode and helical conformation in relation to) L22 ANSWER 19 OF 23 CAPLUS COPYRIGHT 2000 ACS 1969:74287 CAPLUS AN70:74287 DN Adsorption of DNA in the mercury-electrolyte interface TIFlemming, Joachim ΑU Deut. Akad. Wiss. Berlin, Jena, E. Ger. CS Biopolymers (1968), 6(12), 1697-703 SO CODEN: BIPMAA Journal \mathtt{DT} German LA2 (General Biochemistry) CC The adsorption of DNA in the Hg-electrolyte interface has been AB investigated. The effect of this adsorption on the differential capacity of the elec. double layer between a polarized Hg surface and a 0.15M NaCl

soln. contg. DNA was measured by means of the alternating current polarography (Breyer polarograph). The effective a.c. Inder actual conditions (adsorption processes only, small electrolytic resistance, small a.c. frequency, and a.c. amplitude) is directly proportional to the differential double layer capacity. The combination of this method with the application of a stationary Hg drop electrode allows the coverage of the electrode to be followed continuously in the range 0.2 sec. to .apprx.60 sec. diffusion is the rate-controlled step of the adsorption kinetics. Therefore the lowering of the a.c. by the adsorbed DNA is proportional to the surface concn. for partly covered surfaces and reaches a const. value after the surface becomes fully covered. Adsorption of further layers does not affect the differential capacity. This makes it possible to det. the max. surface concn. of the DNA. For that it is necessary to det. the diffusion coeff. of DNA. surface concns. of the native DNA and the relative surface concns. of the denatured DNA in dependence on the potential of the polarized Hg surface were estd. Both surface concns. show a pronounced dependence on the potential with a min. of the surface concn. around -0.4 v. with respect to the normal calomel electrode. This property may be caused by the structure of the adsorption layer depending on the potential. That means that only several segments of the rigid DNA mols. are adsorbed and the other ones remain in the soln. near the surface. The adsorption in the neighborhood of the electrocapillary zero potential at -0.4 v. is strongest, and therefore the fraction of the adsorbed segments has a max. At these potentials consequently, the max. coverage is already reached at relatively low surface concns. DNA adsorption Hg electrode; mercury electrode \mathtt{ST} adsorption DNA Nucleic acids, deoxyribo-ΙŢ RL: PEP (Physical, engineering or chemical process); PROC (Process) (adsorption of, in mercury-electrolyte interface in polarography) L22 ANSWER 20 OF 23 CAPLUS COPYRIGHT 2000 ACS 1971:60984 CAPLUS AN74:60984 DN Adsorption of DNA in the mercury-electrolyte interface TIFlemming, Joachim AU Inst. Mikrobiol. Exptl. Ther., Dtsch. Akad. Wiss. Berlin, Jena, Ger. CS Stud. Biophys. (1968), 8, 209-12 SO CODEN: STBIBN Journal LAGerman 2 (General Biochemistry) CC The adsorption of DNA at mercury-electrolyte interfaces has been AB investigated by means of alternating current polarography. The structure of the adsorption layer depends on the potential of the interface. The adsorption denaturation of the DNA in this interface as supposed by Miller (1961) could not be confirmed. DNA mercury electrode; mercury DNA STelectrode; electrode DNA mercury Nucleic acids, deoxyribo-ITRL: PEP (Physical, engineering or chemical process); PROC (Process) (adsorption of, at mercury-electrolyte interface in polarography) L22 ANSWER 21 OF 23 CAPLUS COPYRIGHT 2000 ACS 1968:464113 CAPLUS AN69:64113 DN Alternating current polarography of nucleosides TIVetterl, Vladimir AU

Ceskoslov. Akad. Ved, Brno, Czech.

CS

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J. Electroanal. Chem. Interfacial Electrochem. (1968), 19(1/2), 169-73
SO
     CODEN: JEIEBC
    Journal
\mathtt{DT}
    English
LA
    77 (Electrochemistry)
CC
     The a.c. polarography of nucleosides currently occurring in
AB
     nucleic acids was studied by using the method of V.
     Vetterl (1966) for measuring the differential capacity of the
     electrode double-layer. The potentials were measured relative to
     the S.C.E. and the concn.-dependence of the shapes of the a.c.
polarograms
     is presented. The a.c. polarograms of nucleosides currently occurring in
     nucleic acids exhibit a min. at .apprx.-0.4 v., caused
     by the adsorption of nucleosides on the electrode surface. At
     higher concns. of deoxycytidine (I), adenosine (II), guanosine, and
     deoxyguanosine, assocn. of the adsorbed mols. occurs in the vicinity of
     -0.4 \ \text{v.} With deoxyadenosine, assocn. of the mol. occurs at .apprx.-1.2
v.
     and with II, at both -0.4 and -1.2 v. As with bases, the transition from
     the nonassocd. to the assocd. state occurs over a closed concn. interval
     in which the adsorption isotherm has an inflection point. With uridine,
     thymidine, and cytidine (III), no assocn. of the adsorbed mols. was
     observed even at concns. approaching satn. value. At pH 7.0, most of the
     nucleosides studied were polarographically nonreducible and the max.
     observed on the a.c. polarograms are of a capacitive character. Only the
     peak for III and I at -1.6 v. is caused by a redn. of cytosine. 20
     references.
     polarog ac nucleosides; nucleosides ac polarog
ST
     Guanosine
IT
     RL: PROC (Process)
        (polarography of, a.c.)
                                               58-96-8 65-46-3
                         58-61-7, reactions
                                                                   951-77-9
     50-89-5, reactions
{	t IT}
     961-07-9
     RL: RCT (Reactant)
        (polarography of, a.c.)
L22 ANSWER 22 OF 23 CAPLUS COPYRIGHT 2000 ACS
     1967:73136 CAPLUS
AN
DN
     66:73136
     Alternating-current polarographic criteria of
TI
     nucleic acid denaturation
     Berg, Hermann; Baer, Horst; Gollmick, F. A.
ΑU
     Deut. Akad. Wiss., Berlin, Ger.
CS
     Biopolymers (1967), 5(1), 61-8
SO
     CODEN: BIPMAA
     Journal
DT
     German
LA
     6 (Biochemical Methods)
CC
     Electrochem. analyses of high-mol.-wt. nucleic acids
AB
     are restricted to the detn. of the adsorption behavior. A.c.
polarography
      (Breyer polarography) can be used for characterizing changes in the
      secondary structure of DNA. The polarogram shows the a.c. of
     the dropping electrode in dependence of the potential which
      ranged 0-2 v. neg. relative to the normal calomel electrode. By
      addn. of native DNA to the supporting electrolyte, the current
      drops in the range of absorption between 0 and 1 v. At 1.16 v.,
      desorption takes place and is indicated by the appearance of a broad
      desorption peak. Denaturation of the double helix causes a sharp
      desorption peak at neg. potentials of the a.c. polarogram.
      criterion for the helix-coil transition is due to the formation of
      unpaired bases which undergo a specific absorption within a narrow
      potential range. In the alk. range, the sharp peak increases and reaches
      its max. at pH >12. In the acid range, no sharp peak is found and the
      broad desorption peak decreases. The best way of following
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conformational

changes is, therefore, to measure the current difference between the curves of the solp with and without DNA at electronapillary zero potential. I eover, the scission of the mol. y ultrasonic action can be followed by the increase of the broad peak of DNA in the absence of any sharp peak. POLAROG DNA DENATURATION; DNA DENATURATION POLAROG; STDENATURATION DNA POLAROG Polarography IT(alternating-current, in structure(secondary) studies) Nucleic acids, deoxyribo-IT RL: PRP (Properties) (structure of, helix-coil transition in, detection by alternating- current polarography) L22 ANSWER 23 OF 23 CAPLUS COPYRIGHT 2000 ACS 1967:479518 CAPLUS AN67:79518 DN Adsorption behavior of nucleic acids from current-time ΤI curves and alternating current polarograms Flemming, Joachim; Berg, Hermann ΑU Deut. Akad. Wiss., Berlin, Ger. CS Abh. Dtsch. Akad. Wiss. Berlin, Kl. Med. (1966), (4), 559-63 SO CODEN: ADWMAX Journal \mathtt{DT} German LA6 (Biochemical Methods) CC The adsorption of nucleic acids at a dropping-Hg AΒ electrode was investigated by measuring the effect of the nucleic acid on the polarographic current-time curves of Cu-EDTA depolarizer and on Breyer alternating current polarograms. The adsorption of RNA and calf thymus DNA was diffusion controlled. The time for complete coverage of the Hg droplet with nucleic acid was obtained from the current-time curves for RNA, but not for DNA, because the overall electrode reaction of the depolarizer was inhibited too weakly. The course of thermal or photochem. denaturation of DNA could then be followed. POLAROG DNA; DNA POLAROG; RNA POLAROG; BERG st

H; FLEMMING J

IT

Nucleic acids, deoxyribo-

(polarography of)

Nucleic acids, ribo-

RL: PROC (Process)

ANSWER 1 OF 16 BIOSIS COPYRIGHT 2000 BIOSIS 1993:24620 BIOSIS AN PREV199395012820 DN Quantification of fluorescence in situ hybridization signals by image ΤI cytometry. Nederlof, P. M.; Van Der Flier, S.; Verwoerd, N. P.; Vrolijk, J.; Raap, AU Α. K. (1); Tanke, H. J. (1) Sylvius Lab., Dep. Ctyochem. Cytometry, Univ. Leiden, Wassenaarseweg CS 72, 2333 Al Leiden Netherlands Cytometry, (1992) Vol. 13, No. 8, pp. 846-852. SO ISSN: 0196-4763. Article DTEnglish LAIn this study we aimed at the development of a cytometric system for AB quantification of specific DNA sequences using fluorescence in situ hybridization (ISH) and digital imaging microscopy. The cytochemical and cytometric aspects of a quantitative ISH procedure were investigated, using human peripheral blood lymphocyte interphase nuclei and probes detecting high copy number target sequences as a model system. These chromosome-specific probes were labeled with biotin, digoxigenin, or fluorescein. The instrumentation requirements are evaluated. Quantification of the fluorescence ISH signals was performed using an epi-fluorescence microscope with a multi-wavelength illuminator, equipped with a cooled charge coupled device (CCD) camera. The performance of the system was evaluated using fluorescing beads and a homogeneosuly fluorescing specimen. Specific image analysis programs were developed for the automated segmentation and analysis of the images provided by ISH. Non-uniform background fluorescence of the nuclei introduces problems in the image analysis segmentation procedures. Different procedures were tested. Up to 95% of the hybridization signals could be correctly segmented using digital filtering techniques (min-max filter) to estimate local background intensities. The choice of the objective lens used for the collection of images was found to be extremely important. High magnification objectives with high numerical aperture, which are frequently used for visualization of fluorescence, are not optimal, since they do not have a sufficient depth of field. The system described was used for quantification of ISH signals and allowed accurate measurement of fluorescence spot intensities, as well as of fluorescence ratios obtained with double-labeled probes. Microscopy Techniques - Cytology and Cytochemistry *01054 CC Cytology and Cytochemistry - Human *02508 Genetics and Cytogenetics - Human *03508 Biochemical Studies - Nucleic Acids, Purines and Pyrimidines Blood, Blood-Forming Organs and Body Fluids - Lymphatic Tissue and Reticuloendothelial System *15008 Hominidae *86215 ВC Major Concepts ITBlood and Lymphatics (Transport and Circulation); Cell Biology; Genetics; Methods and Techniques Miscellaneous Descriptors ITDNA CONTENT; HUMAN PERIPHERAL BLOOD LYMPHOCYTE ORGN Super Taxa Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name Hominidae (Hominidae) ORGN Organism Superterms animals; chordates; humans; mammals; primates; vertebrates

L30 ANSWER 2 OF 16 BLOSIS COPYRIGHT 2000 BIOSIS

AN 1992:497252 BIOS

DN BA94:115777

TI BEHAVIOR OF PERIOD-ALTERED CIRCADIAN RHYTHM MUTANTS OF DROSOPHILA IN LIGHT

DARK CYCLES DIPTERA DROSOPHILIDAE.

AU HAMBLEN-COYLE M J; WHEELER D A; RUTILA J E; ROSBASH M; HALL J C

CS 235 BASSINE BUILD., BRANDEIS UNIV., WALTHAM, MASS. 02254-9110.

SO J INSECT BEHAV, (1992) 5 (4), 417-446. CODEN: JIBEE8.

FS BA; OLD

LA English

AB Adults of Drosophila melanogaster had their locomotor activity monitored under conditions of cycling light and dark (12 h each per cycle). The elementary behavior of wild-type flies under these "LD" conditions fluctuated between levels of high and levels of low activity. Two high-activity peaks occurred within a given cycle: one at about dawn; the other, at around dusk. Such accentuated activity levels gradually subsided

to troughs in the middle of the day and of the night, after which the flies anticipated the next environmental transition by gradually become more active. Descriptions of these activity profiles were augmented by newly developed formal analyses of the "diel rhythm" phases (based in

part

on digital filterings of the raw behavioral data). The applications of these analyses led to objective, automated determination of when in the morning and the evening the flies' activity peaks occur. This normal diel behavior was compared to the locomotor activity and

phase

about

determinations for a series of rhythm variants. Most of these involved mutations at the period (per) locus and germ-line transformants bearing normal or altered forms of **DNA** cloned from this "clock gene." Such genetic variants have been shown previously to exhibit, in constant darkness, strain-specific circadian periods ranging from about 19 to

29 h. We now show that the phases of the evening peaks of activity under LD conditions were correspondingly earlier than normal for the short-period mutants and later than normal for those with long circadian cycle durations. The morning peaks, however, moved (in comparison to the normal phas position) minimally under the influence of a given per variant.

CC Genetics and Cytogenetics - Animal *03506
Behavioral Biology - Animal Behavior *07003
Circadian Rhythms and Other Periodic Cycles *07200
External Effects - Light and Darkness 10604
Movement 12100
Invertebrata, Comparative and Experimental Morphology, Physiology and Pathology - Insecta - Physiology *64076

BC Diptera 75314

IT Miscellaneous Descriptors

DROSOPHILA-MELANOGASTER LOCOMOTOR ACTIVITY CLOCK GENE PHASE ANALYSIS

L30 ANSWER 3 OF 16 BIOSIS COPYRIGHT 2000 BIOSIS

AN 1980:149234 BIOSIS

DN BA69:24230

TI COMPUTER CONTROLLED DOUBLE BEAM SCANNING MICRO SPECTROPHOTOMETRY FOR RAPID

MICROSCOPIC IMAGE RECONSTRUCTIONS.

AU DUCERA P; DE RIBAUPIERRE Y; DE RIAUPIERR F

CS INST. PHYSIOL. UNIV., RUE DE BUGNON 7, CH-01 LAUSANNE, SWITZ.

SO J MICROSC (OXF), (1979) 116 (2), 173-184. CODEN: JMICAR. ISSN: 0022-2720.

FS BA; OLD

LA English

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entire microscopic preparation and their quantita ve evaluation is
    described. Its approcation to the study of neuronal connections is
    discussed in some detail. Brain sections are scanned using a
    computer-controlled microscope for reflectance, fluorescences or
    absorbance signals. Two illuminating beams are used, 1 being amplitude
    modulated. By synchronous detection the 2 signals are recorded
    simultaneosly: e.g., in an autoradiograph, the reflectance (measuring the
    density of the Ag grains in the emulsion) and the absorbance (allowing to
    localize the underlying counterstained cells). The data are stored in a
    computer. Various off-line processing schemes allow the reconstruction of
    the data with respect to the corresponding spatial coordinates.
    Pseudo-3-dimensional, analog or digital, graphic displays may be obtained
    in which the patterns of neuronal connections can be recognized and
    interpreted. A method for the detection of weakly labeled nerve fibers
    based on digital filtering is presented. The whole
    processing for a frontal section of the mouse brain ( 7 .times. 10 nm
     area) takes less than 1 h. In addition to the evaluation of
    microscopically labelled material (grains of autoradographs, horseradish
     peroxidase, nucleic acids) the technique was
     successfully used for the study of naturally fluorescent intracellular
     components in living tissue cultures.
    General Biology - Information, Documentation, Retrieval and Computer
CC
    Applications *00530
    Methods, Materials and Apparatus, General - Photography *01012
    Microscopy Techniques - General and Special Techniques *01052
    Microscopy Techniques - Cytology and Cytochemistry 01054
     Cytology and Cytochemistry - Animal 02506
     Mathematical Biology and Statistical Methods 04500
     Radiation - Radiation and Isotope Techniques 06504
     Biochemical Studies - Nucleic Acids, Purines and Pyrimidines 10062
     Biochemical Studies - Proteins, Peptides and Amino Acids 10064
     Biochemical Studies - Porphyrins and Bile Pigments 10065
     Biochemical Studies - Minerals 10069
     Biophysics - General Biophysical Techniques 10504
     Biophysics - Biocybernetics
                                 10515
     Enzymes - Methods 10804
     Enzymes - Physiological Studies 10808
     Anatomy and Histology, General and Comparative - Microscopic and
     Ultramicroscopic Anatomy *11108
     Nervous System - General; Methods 20501
     Nervous System - Physiology and Biochemistry 20504
     Tissue Culture, Apparatus, Methods and Media 32500
     Plant Physiology, Biochemistry and Biophysics - Enzymes 51518
     Cruciferae 25880
BC
     Muridae 86375
     Miscellaneous Descriptors
IT
        MOUSE BRAIN AUTO RADIOGRAPHY REFLECTANCE ABSORBANCE FLUORESCENCE
TISSUE
        CULTURE HORSERADISH PEROXIDASE
     9003-99-0 (PEROXIDASE)
RN
    ANSWER 4 OF 16 MEDLINE
L30
     93092792
                  MEDLINE
AN
     93092792
DN
     Quantification of fluorescence in situ hybridization signals by image
TI
     cytometry.
     Nederlof P M; van der Flier S; Verwoerd N P; Vrolijk J; Raap A K; Tanke H
ΑU
     Sylvius Laboratory, Department of Cytochemistry and Cytometry, University
CS
     of Leiden, The Netherlands..
     CYTOMETRY, (1992) 13 (8) 846-52.
SO
     Journal code: D92. ISSN: 0196-4763.
CY
     United States
     Journal; Article; (JOURNAL ARTICLE)
\mathsf{DT}
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A method for the automated collection of various specific data from an

AΒ

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English
LA
     Priority Journals
FS
EM
     199303
     In this study we aimed at the development of a cytometric system for
AB
     quantification of specific DNA sequences using fluorescence in
     situ hybridization (ISH) and digital imaging microscopy. The cytochemical
     and cytometric aspects of a quantitative ISH procedure were investigated,
     using human peripheral blood lymphocyte interphase nuclei and probes
     detecting high copy number target sequences as a model system. These
     chromosome-specific probes were labeled with biotin, digoxigenin, or
     fluorescein. The instrumentation requirements are evaluated.
     Quantification of the fluorescence ISH signals was performed using an
     epi-fluorescence microscope with a multi-wavelength illuminator, equipped
     with a cooled charge couple device (CCD) camera. The performance of the
     system was evaluated using fluorescing beads and a homogeneously
     fluorescing specimen. Specific image analysis programs were developed for
     the automated segmentation and analysis of the images provided by ISH.
     Non-uniform background fluorescence of the nuclei introduces problems in
     the image analysis segmentation procedures. Different procedures were
     tested. Up to 95% of the hybridization signals could be correctly
     segmented using digital filtering techniques (min-max
     filter) to estimate local background intensities. The choice of the
     objective lens used for the collection of images was found to be
extremely
     important. High magnification objectives with high numerical aperture,
     which are frequently used for visualization of fluorescence, are not
     optimal, since they do not have a sufficient depth of field. The system
     described was used for quantification of ISH signals and allowed accurate
     measurement of fluorescence spot intensities, as well as of fluorescence
     ratios obtained with double-labeled probes.
     Check Tags: Human; Support, Non-U.S. Gov't
CT
      Analog-Digital Conversion
     *Cell Nucleus: UL, ultrastructure
      Chromosomes, Human, Pair 1
      Chromosomes, Human, Pair 7
     *DNA: AN, analysis
      DNA Probes
      DNA, Satellite: AN, analysis
      Image Processing, Computer-Assisted: IS, instrumentation
     *Image Processing, Computer-Assisted: MT, methods
      In Situ Hybridization, Fluorescence: IS, instrumentation
     *In Situ Hybridization, Fluorescence: MT, methods
      Interphase
     *Lymphocytes: UL, ultrastructure
      Microscopy, Fluorescence: IS, instrumentation
      Photomicrography: IS, instrumentation
     9007-49-2 (DNA)
RN
     0 (DNA Probes); 0 (DNA, Satellite)
CN
    ANSWER 5 OF 16 MEDLINE
L30
     81096628
                  MEDLINE
AN
     81096628
DN
     Computer-controlled double-beam scanning microspectrophotometry for rapid
TI
     microscopic image reconstructions.
     Kucera P; de Ribaupierre Y; de Ribaupierre F
ΑU
     JOURNAL OF MICROSCOPY, (1979 Jul) 116 (2) 173-84.
SO
     Journal code: J5V. ISSN: 0022-2720.
     ENGLAND: United Kingdom
CY
     Journal; Article; (JOURNAL ARTICLE)
DT
     English
LA
     Priority Journals
FS
     198105
EM
     A method for automated collection of various specific data from an entire
AB
     microscopical preparation and their quantitative evaluation is described.
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Its application to the study of neuronal connections is discussed in some

detail. Brain sections are scanned using a computer-controlled microscope for reflectance, forescences or absorbance signal Two illuminating beams are used, on of them being amplitude modulated. By means of a synchronous detection the two signals are recorded simultaneously: for example, in an autoradiograph, the reflectance (measuring the density of the silver grains in emulsion) and the absorbance (allowing to localize the underlying counterstained cells). The data are stored in a computer. Various off-line processing schemes allow the reconstruction of the data with respect to the corresponding spatial coordinates. Thus pseudo-three-dimensional, analogue or digital, graphic displays may be obtained in which the patterns of neuronal connections can be recognized and interpreted. A method for the detection of weakly labelled nerve fibres based on digital filtering is presented. The whole processing for a frontal section of the mouse brain (7 X 10 mm area) takes less than 1 h. In addition to the evaluation of microscopically labelled material (grains of autoradiographs, horseradish peroxidase, nucleic acids) the technique described has been successfully used for the study of naturally fluorescent intracellular components in living tissue cultures. Check Tags: Animal; Support, Non-U.S. Gov't Autoradiography Brain: CY, cytology Chick Embryo Computers Mice *Microscopy: MT, methods *Neural Pathways Neurons: CY, cytology *Spectrophotometry Staining L30 ANSWER 6 OF 16 CAPLUS COPYRIGHT 2000 ACS 1999:365091 CAPLUS AN 131:196538 DN Analysis of ultrasensitive fluorescence experiments TISun, Yuxing; Whitehead, Bruce A.; Davis, Lloyd M. ΑU Center for Laser Applications, Univ. of Tennessee Space Institute, CS Tullahoma, TN, USA Proc. SPIE-Int. Soc. Opt. Eng. (1999), 3602 (Advances in Fluorescence SO Sensing Technology IV), 379-390 CODEN: PSISDG; ISSN: 0277-786X SPIE-The International Society for Optical Engineering PΒ Journal English LA9-5 (Biochemical Methods) CCSection cross-reference(s): 3 DNA sequencing and several other applications of single- mol. AB detection (SMD) currently under development utilize spectroscopic measurements for categorization of different types of fluorophores. In the collection and anal. of data from such expts., the photon signals are sorted into different channels, depending upon their arrival time, emission wavelength, or other distinguishable properties. If the photon statistics are adequate, max.-likelihood estn. (MLE) techniques can be successfully applied to det. which fluorophore is present. However, data anal. using neural network (NN) methods can offer several advantages. We consider data from a Monte Carlo simulation of SMD in a flow-cell, in which a time-resolved fluorescence decay profile is accumulated for each photon burst. A 2-layer NN, with sigmoid as the activation function, is trained on a set of simulated data using back-propagation and the (delta) - learning rule, and then used for identification of photon bursts in subsequent simulations. The NN is able to consider addnl. input

parameters, such as the amplitudes of the weighted-sliding-sum

durations of the bursts. It can yield superior identification of photon

digital-filter output of the photon bursts and the

CT

DT

bursts, particularly in cases where the fluorophores have disparate fluorescence quanta efficiencies, absorption cross ections, or photodegrdn. efficiencies, or where the categorization includes other possibilities, such as background fluctuations, or the simultaneous presence of both fluorophores. neural network fluorescence single mol detection DNA sequencing Fluorometry (max.-likelihood estn. and neural network methods for anal. of fluorescence single- mol. detection) DNA sequence analysis Mathematical methods (max.-likelihood estn. and neural network methods for anal. of fluorescence single- mol. detection in) Simulation and Modeling, physicochemical (neural network; max.-likelihood estn. and neural network methods for anal. of fluorescence single- mol. detection in) RE.CNT 22 (1) Bunfield, D; Appl Opt 1995, V34, P3208 (2) Bunfield, D; Thesis University of Tennessee 1997 (3) Davis, L; BiOS Europe Conference 1998, P282 (4) Davis, L; Biomedical Sensors Fibers and Optical Delivery Systems 1999 (5) Davis, L; Book of Abstracts (6) Davis, L; SPIE Proceedings V3570 (7) Davis, L; The Fifth International Conference on Methods and Applications Fluorescence Spectroscopy 1997, P27 (8) Dorre, K; Bioimaging 1997, V5, P139 CAPLUS (9) Enderlein, J; Chem Phys Lett 1997, V270, P464 CAPLUS (10) Kollner, M; Appl Opt 1993, V32, P806 (11) Kollner, M; Chem Phys Lett 1992, V200, P199 (12) Krose, B; An Introduction to Neural networks 8th ed 1996 (13) Li, L; Appl Opt 1993, V32, P806 (14) Lieberwirth, U; Anal Chem 1998, V70, P4771 CAPLUS (15) Sauer, M; Applied Fluorescence in Chemistry Biology and Medicine 1999 (16) Sauer, M; Bioimaging 1998, V6, P14 CAPLUS (17) Soper, S; J Opt Soc Am B 1992, V9, P1761 CAPLUS (18) Soper, S; Photochem and Photobiol 1993, V57, P972 CAPLUS (19) Werner, J; Advances in Fluorescence Sensing Technology 1999, V4 (20) Werner, J; BiOS Conference 1999 (21) Werner, J; paper 40 in SPIE Proceedings V3602 (22) Zander, C; Appl Phys B 1996, V63, P517 L30 ANSWER 7 OF 16 CAPLUS COPYRIGHT 2000 ACS AN 1999:228827 CAPLUS 131:69253 Computer simulation of gene detection without PCR by single molecule detection Davis, Lloyd M.; Williams, John G. K.; Lamb, Don T. Center for Laser Applications, University of Tennessee Space Institute, Tullahoma, TN, 37388, USA Proc. SPIE-Int. Soc. Opt. Eng. (1999), 3570 (Biomedical Sensors, Fibers, and Optical Delivery Systems), 282-293 CODEN: PSISDG; ISSN: 0277-786X SPIE-The International Society for Optical Engineering Journal English 3-6 (Biochemical Genetics) Section cross-reference(s): 9 Pioneer Hi-Bred is developing a low-cost method for rapid screening of DNA, for use in research on elite crop seed genetics. Unamplified

genomic DNA with the requisite base sequence is simultaneously

labeled by two different colored fluorescent probes, which hybridize near

the selected gene. Dual-channel single mol. detection (SMD) within a

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cell then provides a sensitive and specific assay for the gene. technique has been emonstrated using frequency-doved Nd:YAG laser excitation of two sible-wavelength dyes. A proto pe instrument employing IR fluorophores and laser diodes for excitation has been developed. Here, we report results from a Monte Carlo simulation of the new instrument, in which exptl. detd. photophys. parameters for candidate IR dyes are used for parametric studies of exptl. operating conditions. Our findings demonstrate the feasibility of the approach for selected fluorophores, and identify suitable operating conditions. Fluorophore photostability is found to be a key factor in detg. the instrument sensitivity. Most IR dyes have poor photostability, resulting in inefficient SMD. However, the normalized cross-correlation function of the photon signals from each of the two channels can still yield a discernable peak, provided that the concn. of dual-labeled mols. is sufficiently high. Further, for low concns., processing of the two photon streams with Gaussian weighted sliding sum digital filters and selection of simultaneously occurring peaks can also provide a sensitive indicator of the presence of dual-labeled mols., although accidental coincidences must be considered in the interpretation of results. computer simulation gene screening single mol detection Nucleic acid hybridization (DNA-DNA; computer simulation of gene detection without PCR by single mol. detection) Dyes (IR; computer simulation of gene detection without PCR by single mol. detection) Simulation and Modeling, physicochemical (Monte Carlo; computer simulation of gene detection without PCR by single mol. detection) Fluorescence Molecules (computer simulation of gene detection without PCR by single mol. detection) Gene RL: BSU (Biological study, unclassified); BIOL (Biological study) (detection of; computer simulation of gene detection without PCR by single mol. detection) Genetic methods (dual-channel single mol. detection (SMD); computer simulation of gene detection without PCR by single mol. detection) Fluorescent substances (photostability of, key factor in detg. the instrument sensitivity; computer simulation of gene detection without PCR by single mol. detection) Probes (nucleic acid) RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (two different colored fluorescent; computer simulation of gene detection without PCR by single mol. detection)

ΙT

Fluorescent dyes IT

(two probes labeled with different; computer simulation of gene detection without PCR by single mol. detection)

ITDNA

RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study)

(unamplified genomic, gene detection in; computer simulation of gene detection without PCR by single mol. detection)

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- (1) Bunfield, D; Appl Opt 1998, V37, P2315 CAPLUS
- (2) Castro, A; Anal Chem 1997, V69, P3915 CAPLUS
- (3) Loudon, R; The Quantum Theory of Light 1st ed 1973, P210
- (4) Lundgren, T; J Basic Eng 1964, V86, P620
- (5) Soper, S; Photochem and Photobiol 1993, V57, P972 CAPLUS

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ANSWER 8 OF 16 CAMBUS COPYRIGHT 2000 ACS
L30
     1996:304534 CAPLU
AN
     125:29407
DN
     Single molecule fluorescence burst detection of DNA separated by
TI
     capillary electrophoresis
     Haab, Brian B.; Mathies, Richard A.
AU
     Department of Chemistry, University of California, Berkeley, CA, 94720,
CS
     USA
     Proc. SPIE-Int. Soc. Opt. Eng. (1996), 2705(Fluorescence Detection IV),
SO
     162-169
     CODEN: PSISDG; ISSN: 0277-786X
     Journal
\mathtt{DT}
    English
LA
     9-5 (Biochemical Methods)
CC
     A method has been developed for detecting DNA sepd. by capillary
AB
     gel electrophoresis using single mol. photon burst counting. A confocal
     fluorescence microscope was used to observe the fluorescence bursts from
     single mols. of DNA multiply labeled with a thiazole orange
     deriv. as they passed through the .apprx.2 .mu.m diam. focused laser
beam.
     Amplified photoelectron pulses from the photomultiplier are grouped into
     bins of from 360-450 .mu.s in duration, and the resulting histogram
stored
     in a computer for anal. Solns. of M13 DNA were first flowed
     through the capillary at various concns., and the resulting data were
used
     to optimize the parameters for digital filtering using
     a low-pass Fourier filter, selecting a discriminator level for peak
     detection, and applying a peak-calling algorithm. The optimized single
     mol. counting method was then used to detect a sepn. of pBR 322
     DNA from pRL 277 DNA. Clusters of discrete fluorescence
     bursts were obsd. at the expected appearance time of each DNA
     band. These sepns. were easily detected when only 50 to 100 mols. of
     DNA per band traveled through the detection region. This new
     detection technol. should lead to the routine anal. of DNA in
     capillary columns with an on-column sensitivity of .apprx. 100 DNA
     mols. per band or better.
     DNA detection single mol fluorescence burst
ST
     Photon
IT
        (single mol. fluorescence burst detection of DNA sepd. by
        capillary electrophoresis)
     Deoxyribonucleic acids
IT
     RL: ANT (Analyte); ANST (Analytical study)
        (single mol. fluorescence burst detection of DNA sepd. by
        capillary electrophoresis)
L30 ANSWER 9 OF 16 CAPLUS COPYRIGHT 2000 ACS
     1995:800379 CAPLUS
AN
     123:247800
DN
     Single molecule fluorescence burst detection of DNA fragments
TI
     separated by capillary electrophoresis
     Haab, Brian B.; Mathies, Richard A.
AU
     Department of Chemistry, University of California, Berkeley, CA, 94720,
CS
     USA
     Anal. Chem. (1995), 67(18), 3253-60
SO
     CODEN: ANCHAM; ISSN: 0003-2700
     Journal
\mathsf{DT}
     English
LA
     3-1 (Biochemical Genetics)
CC
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A method has been developed for detecting DNA sepd. by capillary

bursts from single mols. of DNA multiply labeled with the

gel electrophoresis (CGE) using single mol. photon burst counting. A confocal fluorescence microscope was used to observe the fluorescence

Section cross-reference(s): 9

AB

thiazole orange deriv. TO6 as they passed through the .apprx.2-.mu.m diam. focused laser beam Amplified photoelectron pulses from the photomultiplier are grouped into bins of 360-450 .mu.s in duration, and the resulting histogram is stored in a computer for anal. Solns. of M13 DNA were first flowed through the capillary at various concns., and the resulting data were used to optimize the parameters for digital filtering using a low-pass Fourier filter, selecting a discriminator level for peak detection, and applying a peak-calling algorithm. Statistical analyses showed that (i) the no. of M13 mols. counted vs. concn. was linear with slope = 1, (ii) the av. burst duration was consistent with the expected transit time of a single mol. through the laser beam, and (iii) the no. of detected mols. was consistent with single mol. detection. The optimized single mol. counting method was then applied to an electrophoretic sepn. of M13 DNA and to a sepn. of pBR322 DNA from pRL277 DNA. Clusters of discreet fluorescence bursts were obsd. at the expected appearance time of each DNA band. The autocorrelation function of these data indicated transit times that were consistent with the obsd. electrophoretic velocity. These sepns. were easily detected when only 50-100 mols. of DNA per band traveled through the detection region. This new detection technol. should lead to the routine anal. of DNA in capillary columns with an on-column sensitivity of .apprx.100 DNA mols./band or better. DNA capillary electrophoresis single mol fluorescence ST Lasers IT(a confocal fluorescence microscope was used to observe the fluorescence bursts from single mols. of DNA multiply labeled with the thiazole orange deriv. TO6 as they passed through the .apprx.2-.mu.m diam. focused laser beam) ITPhoton (a method has been developed for detecting DNA sepd. by capillary gel electrophoresis using single mol. photon burst counting) Plasmid and Episome IT(pRL277; the optimized single mol. fluorescence burst detection of DNA fragments sepd. by capillary electrophoresis was then applied to an electrophoretic sepn. of M13 DNA and to a sepn. of pBR322 DNA from pRL277 DNA) Fluorescence ΙT (single mol. fluorescence burst detection of DNA fragments sepd. by capillary electrophoresis) Deoxyribonucleic acids IT RL: ANT (Analyte); ANST (Analytical study) (the optimized single mol. fluorescence burst detection of DNA fragments sepd. by capillary electrophoresis was then applied to an electrophoretic sepn. of M13 DNA and to a sepn. of pBR322 DNA from pRL277 DNA) Virus, bacterial IT(M13, the optimized single mol. fluorescence burst detection of DNA fragments sepd. by capillary electrophoresis was then applied to an electrophoretic sepn. of M13 DNA and to a sepn. of pBR322 DNA from pRL277 DNA) Electrophoresis and Ionophoresis IT(gel, capillary, single mol. fluorescence burst detection of DNA fragments sepd. by capillary electrophoresis) Plasmid and Episome IT(pBR322, the optimized single mol. fluorescence burst detection of DNA fragments sepd. by capillary electrophoresis was then applied to an electrophoretic sepn. of M13 DNA and to a sepn. of pBR322 DNA from pRL277 DNA) 153087-66-2, TO 6 IT

RL: BAC (Biological activity or effector, except adverse); BUU (Biological use, unclassified) BIOL (Biological study); USES ((a confocal fluorescence microscope was used to observe the fluorescence bursts from single mols. of DNA multiply labeled with the thiazole orange deriv. TO6 as they passed through the .apprx.2-.mu.m diam. focused laser beam) L30 ANSWER 10 OF 16 EMBASE COPYRIGHT 2000 ELSEVIER SCI. B.V. 1998166655 EMBASE ANVisualization of single RNA transcripts in situ. TIFemino A.M.; Fay F.S.; Fogarty K.; Singer R.H. AU R.H. Singer, Department of Anatomy, Albert Einstein College of Medicine, CS Bronx, NY 10461, United States Science, (24 Apr 1998) 280/5363 (585-590). SO Refs: 24 ISSN: 0036-8075 CODEN: SCIEAS CYUnited States Journal; Article DTClinical Biochemistry FS 029 English $_{
m LA}$ \mathtt{SL} English Fluorescence in situ hybridization (FISH) and digital imaging microscopy AB were modified to allow detection of single RNA molecules. Oligodeoxynucleotide probes were synthesized with five fluorochromes per molecule, and the light emitted by a single probe was calibrated. Points single messenger RNA molecules. Analysis of .beta.-actin transcription sites after serum induction revealed synchronous and initiation and termination and messenger RNA processing could be determined by positioning probes along the transcription unit. This approach extends the power of FISH to yield quantitative molecular information on a single cell. Medical Descriptors: CT

of light in exhaustively deconvolved images of hybridized cells gave fluorescent intensities and distances between probes consistent with cyclical transcription from single genes. The rates of transcription

*rna analysis

*rna processing

fluorescence in situ hybridization

digital filtering

infrared radiation

transcription regulation

binding site

dna probe

priority journal

Drug Descriptors:

*messenger rna: EC, endogenous compound

*beta actin: EC, endogenous compound

- L30 ANSWER 11 OF 16 EMBASE COPYRIGHT 2000 ELSEVIER SCI. B.V.
- 92338287 EMBASE AN
- 1992338287 DN
- Quantification of fluorescence in situ hybridization signals by image TIcytometry.
- Nederlof P.M.; Van der Flier S.; Verwoerd N.P.; Vrolijk J.; Raap A.K.; ΑU Tanke H.J.
- Sylvius Laboratory, Dept. of Cytochemistry/Cytometry, University of CS Leiden, Wassenaarseweg 72,2333 AL Leiden, Netherlands
- Cytometry, (1992) 13/8 (846-852). SO ISSN: 0196-4763 CODEN: CYTODQ
- CY United States
- Journal; Article DT
- 022 Human Genetics FS
 - Hematology 025

Immunology, Serology and Transplantation 026 Biophysics Bioengineering and Medical Instantation 027

English LA

English \mathtt{SL} In this study we aimed at the development of a cytometric system for AB quantification of specific DNA sequences using fluorescence in situ hybridization (ISH) and digital imaging microscopy. The cytochemical and cytometric aspects of a quantitative ISH procedure were investigated, using human peripheral blood lymphocyte interphase nuclei and probes detecting high copy number target sequences as a model system. These chromosome-specific probes were labeled with biotin, digoxigenin, or fluorescein. The instrumentation requirements are evaluated. Quantification of the fluorescence ISH signals was performed using an epi-fluorescence microscope with a multi-wave-length illuminator,

equipped

with a cooled charge couple device (CCD) camera. The performance of the system was evaluated using fluorescing beads and a homogeneously fluorescing specimen. Specific image analysis programs were developed for the automated segmentation and analysis of the images provided by ISH. Non-uniform background fluorescence of the nuclei introduces problems in the image analysis segmentation procedures. Different procedures were tested. Up to 95% of the hybridization signals could be correctly segmented using digital filtering techniques (min-max filter) to estimate local background intensities. The choice of the

objective lens used for the collection of images was found to be extremely

important. High magnification objectives with high numerical aperture, which are frequently used for visualization of fluorescence, are not optimal since they do not have a sufficient depth of field. The system described was used for quantification of ISH signals and allowed accurate measurement of fluorescence spot intensities, as well as fluorescence ratios obtained with double-labeled probes.

Medical Descriptors: CT

*dna sequence

- *fluorescence
- *in situ hybridization
- *quantitative assay

adult

article

chromosome 1

human

human cell

image analysis

normal human

priority journal

*dna probe

- ANSWER 12 OF 16 EMBASE COPYRIGHT 2000 ELSEVIER SCI. B.V. L30
- 79258416 EMBASE AN
- 1979258416 DN
- Computer-controlled double-beam scanning microspectrophotometry for rapid TImicroscopic image reconstructions.
- Kucera P.; De Ribaupierre Y.; De Ribaupierre F. UΑ
- Inst. Physiol. Univ., CH-1011 Lausanne, Switzerland CS
- Journal of Microscopy, (1979) 116/2 (173-184). SO
 - CODEN: JMICAR
- United Kingdom CY
- Journal \mathtt{DT}
- Anatomy, Anthropology, Embryology and Histology FS 001 Biophysics, Bioengineering and Medical Instrumentation 027
- English LA
- A method for automated collection of various specific data from an entire ABmicroscopical preparation and their quantitative evaluation is described. Its application to the study of neuronal connections is discussed in some detail. Brain sections are scanned using a computer-controlled microscope

for reflectance, fluorescences or absorbance signals. Two illuminating beams are used, or of them being amplitude modulat By means of a synchronous detect In the two signals are recorded multaneously: for example, in an autoradiograph, the reflectance (measuring the density of the silver grains in the emulsion) and the absorbance (allowing to localize the underlying counterstained cells). The data are stored in a computer. Various off-line processing schemes allow the reconstruction of the data with respect to the corresponding spatial coordinates. Thus pseudo-three-dimensional, analogue or digital, graphic displays may be obtained in which the patterns of neuronal connections can be recognized and interpreted. A method for the detection of weakly labelled nerve fibres based on digital filtering is presented. The whole processing for a frontal section of the mouse brain (7 \times 10 mm area) takes less than 1 hr. In addition to the evaluation of microscopically labelled material (grains of autoradiographs, horseradish peroxidase, nucleic acids) the technique described has been successfully used for the study of naturally fluorescent intracellular components in living tissue cultures. Medical Descriptors: *image *microscopy *microspectrophotometry methodology electron microscopy computer analysis L30 ANSWER 13 OF 16 SCISEARCH COPYRIGHT 2000 ISI (R) 2000:100010 SCISEARCH The Genuine Article (R) Number: 279KM Digitally filtered molecular dynamics: The frequency specific control of molecular dynamics simulations Phillips S C; Essex J W (Reprint); Edge C M UNIV SOUTHAMPTON, DEPT CHEM, SOUTHAMPTON SO17 1BJ, HANTS, ENGLAND (Reprint); UNIV SOUTHAMPTON, DEPT CHEM, SOUTHAMPTON SO17 1BJ, HANTS, ENGLAND; SMITHKLINE BEECHAM PHARMACEUT, HARLOW CM19 5AD, ESSEX, ENGLAND CYA ENGLAND JOURNAL OF CHEMICAL PHYSICS, (8 FEB 2000) Vol. 112, No. 6, pp. 2586-2597. Publisher: AMER INST PHYSICS, CIRCULATION FULFILLMENT DIV, 500 SUNNYSIDE BLVD, WOODBURY, NY 11797-2999. ISSN: 0021-9606. Article; Journal PHYS English Reference Count: 28 REC A new method for modifying the course of a molecular dynamics computer simulation is presented. Digitally filtered molecular dynamics (DFMD) applies the well-established theory of digital filters to molecular dynamics simulations, enabling atomic motion to be enhanced or suppressed in a selective manner solely on the basis of frequency. The basic theory of digital filters and its application to molecular dynamics simulations is presented, together with the application of DFMD to the simple systems of single molecules of water and butane. The extension of the basic theory to the condensed phase is then described followed by its application to liquid phase butane and the Syrian hamster prion protein. The high degree of selectivity and control offered by DFMD, and its ability to enhance the rate of conformational change in butane and in the prion protein, is demonstrated. (C) 2000 American Institute of Physics. [S0021-9606(00)52805-0]. PHYSICS, ATOMIC, MOLECULAR & CHEMICAL

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,	Year	•	PG (RPG)	Referenced Work (RWK) +==========
ASKAR A		100	19165	J PHYS CHEM-US
BECKER O M	•	70	3514	PHYS REV LETT
CASE D A	1997	Ì		AMBER 5
CORNELL W D	1995	1117	5179	J AM CHEM SOC
DARDEN T	11993	98	10089	J CHEM PHYS
DAUBEROSGUTHORPE P	1990	1112	17921	J AM CHEM SOC
DAUBEROSGUTHORPE P	1996	10	177	J COMPUT AID MOL DES
GOLDFARB L G	1992	258	1806	SCIENCE
GREST G S	1980	136	875	SOLID STATE COMMUN
HUBER T	1998	102	5937	J PHYS CHEM A
JORGENSEN W L	1995		[BOSS VERSION 3 6
JORGENSEN W L	1984	106	6638	J AM CHEM SOC
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LEVITT M	11991	1220	11	J MOL BIOL
LU H	1998	75	662	BIOPHYS J
OSGUTHORPE D J	11992	10	178	J MOL GRAPHICS
PAN K M	1993	190	10962	P NATL ACAD SCI USA
PARCHMENT O G				IN PRESS PROTEINS ST
PRESS W H	11992	1		NUMERICAL RECIPES C
RYCKAERT J P	1977	23	1327	J COMPUT PHYS
SAFAR J	1993	12	12206	PROTEIN SCI
SESSIONS R B	1989	210	617	J MOL BIOL
SESSIONS R B	1995	199	19034	J PHYS CHEM-US
SMITH W	11996	114	136	J MOL GRAPHICS
TELEMAN O	1987	160	193	MOL PHYS
WEINER S J	1984	106	1765	J AM CHEM SOC
WILLIAMS C S	1986	1	!	DESIGNING DIGITAL FI
XU D	1995	103	3124	J CHEM PHYS

L30 ANSWER 14 OF 16 SCISEARCH COPYRIGHT 2000 ISI (R)

AN 95:622280 SCISEARCH

GA The Genuine Article (R) Number: RU331

TI SINGLE-MOLECULE FLUORESCENCE BURST DETECTION OF **DNA** FRAGMENTS SEPARATED CAPILLARY ELECTROPHORESIS

AU HAAB B B; MATHIES R A (Reprint)

CS UNIV CALIF BERKELEY, DEPT CHEM, BERKELEY, CA, 94720 (Reprint); UNIV CALIF BERKELEY, DEPT CHEM, BERKELEY, CA, 94720

CYA USA

the

SO ANALYTICAL CHEMISTRY, (15 SEP 1995) Vol. 67, No. 18, pp. 3253-3260. ISSN: 0003-2700.

DT Article; Journal

FS PHYS; LIFE

LA ENGLISH

REC Reference Count: 44

AB A method has been developed for detecting DNA separated by capillary gel electrophoresis (CGE) using single molecule photon burst counting, A confocal fluorescence microscope was used to observe the fluorescence bursts from single molecules of DNA multiply labeled with the thiazole orange derivative TO6 as they passed through

similar to 2-mu m diameter focused laser beam. Amplified photoelectron pulses from the photomultiplier are grouped into bins of 360-450 mu s in duration, and the resulting histogram is stored in a computer for analysis, Solutions of M13 DNA were first flowed through the capillary at various concentrations, and the resulting data were used to optimize the parameters for digital filtering using a

optimize the parameters for **digital filtering** using a lowpass Fourier filter, selecting a discriminator level for peak detection, and applying a peak-calling algorithm, Statistical analyses showed that (i) the number of M13 molecules counted versus concentration

was linear with slope = 1, (ii) the average burst duration was consistent with the expected ansit time of a single molecule hrough the laser beam, and (iii) the humber of detected molecules was consistent with single molecule detection. The optimized single molecule counting method was then applied to an electrophoretic separation of M13 DNA and to a separation of pBR 322 DNA from pRL 277 DNA.

Clusters of discreet fluorescence bursts were observed at the expected appearance time of each DNA band, The autocorrelation function of these data indicated transit times that were consistent with the observed electrophoretic velocity. These separations were easily detected when only 50-100 molecules of DNA per band traveled through the detection region. This new detection technology should lead to the

analysis of **DNA** in capillary columns with an on-column sensitivity of similar to 100 **DNA** molecules/band or better.

CC CHEMISTRY, ANALYTICAL

STP KeyWords Plus (R): LASER-INDUCED FLUORESCENCE; GEL-ELECTROPHORESIS; SPECTROSCOPY; PHYCOERYTHRIN; EXCITATION; MICROSCOPY; SIZE

RF 93-0744 003; PERSISTENT SPECTRAL HOLE-BURNING; SINGLE MOLECULES; HIGH-RESOLUTION SPECTROSCOPY; OPTICALLY DRIVEN QUANTUM NETWORKS;

DISPERSED FLUORESCENCE

93-2117 002; CAPILLARY ELECTROPHORESIS; SIMULTANEOUS CHIRAL SEPARATION; SELECTIVITY MANIPULATION IN MICELLAR ELECTROKINETIC CHROMATOGRAPHY 93-3721 002; PULSED-FIELD GEL-ELECTROPHORESIS; DNA DOUBLE-STRAND BREAKS; YEAST CHROMOSOMES

Referenced Author (RAU)	Year (RPY)		PG (RPG)	Referenced Work (RWK)
AMBROSE W P			7150	J CHEM PHYS
AMBROSE W P	11994	265	364	SCIENCE
BASCHE T	1992	355	335	NATURE
BENSON S C	1993	21	5720	NUCLEIC ACIDS RES
BENSON S C	1993	21	15727	NUCLEIC ACIDS RES
BETZIG E	1993	262	1422	SCIENCE
BLACK T A	11993	19	177	MOL MICROBIOL
CASTRO A	1993	65	849	ANAL CHEM
CLARK S M	1993	1215	163	ANAL BIOCHEM
EWING A G	1989	61	A 292	ANAL CHEM
GLAZER A N	1990	87	3851	P NATL ACAD SCI USA
GOODWIN P M	11993	121	1803	NUCLEIC ACIDS RES
HELL S	1993	169	391	J MICROSC-OXFORD
HIRSCHFELD T	11976	15	12965	APPL OPTICS
HJERTEN S	1985	347	191	J CHROMATOGR
INGLE J D	11988	†	CH 5	SPECTROCHEMICAL ANAL
ISHIKAWA M	11994	33	1571	JPN J APPL PHYS PT 1
LANDERS J P	11993	114	98	BIOTECHNIQUES
LEE Y H	1994	66	4142	ANAL CHEM
MATHIES R A	11990	62	1786	ANAL CHEM
METS U	1994	14	259	J FLUORESC
MOERNER W E	1989	62	2535	PHYS REV LETT
MOERNER W E	1994	•	146	SCIENCE
NGUYEN D C				ANAL CHEM
NIE S M				SCIENCE
ORRIT M				J LUMIN
PECK K	•	-		P NATL ACAD SCI USA
PERKINS T T	11994	1264	819	
PETERSEN N O	1986	49		BIOPHYS J
PRESS W H	11992	ļ		NUMERICAL RECIPES C
SCHAFER D A	1992	•	444	NATURE
SCHWARTZ D C	•	-	520	
SHERA E B			553	
SMITH S B	•			SCIENCE
SMITH S B	1992	258	1122	SCIENCE

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|1991 |63
                                         |ANAL CHEM
                                  | 432
SOPER S A
                                         J OPT SOC AM
                      992 | 9
                                  11761
SOPER S A
                       994 | 369
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                                         NATURE
TRAUTMAN J K
                                        ANAL CHEM
                      11991 | 63
                                  11027
WHITTEN W B
                                  12030
                      |1993 |62
                                         |APPL PHYS LETT
WILKERSON C W
                      |1987 |12
                                  1227
                                         OPT LETT
WILSON T
                                  |11348 | P NATL ACAD SCI USA
                      |1994 |91
WOOLLEY A T
                      |1994 |265
                                  |361
                                         SCIENCE
XIE X S
                      |1994 |66
                                  |1941 |ANAL CHEM
ZHU H P
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L30 ANSWER 15 OF 16 SCISEARCH COPYRIGHT 2000 ISI (R)

AN 92:641565 SCISEARCH

GA The Genuine Article (R) Number: JV644

TI QUANTIFICATION OF FLUORESCENCE INSITU HYBRIDIZATION SIGNALS BY IMAGE CYTOMETRY

AU NEDERLOF P M; VANDERFLIER S; VERWOERD N P; VROLIJK J; RAAP A K (Reprint); TANKE H J

CS LEIDEN UNIV, DEPT CYTOCHEM & CYTOMETRY, SYLVIUS LAB, WASSENAARSEWEG 72, 2333 AL LEIDEN, NETHERLANDS

CYA NETHERLANDS

SO CYTOMETRY, (1992) Vol. 13, No. 8, pp. 846-852. ISSN: 0196-4763.

DT Article; Journal

FS LIFE

LA ENGLISH

REC Reference Count: 23

In this study we aimed at the development of a cytometric system for quantification of specific DNA sequences using fluorescence in situ hybridization (ISH) and digital imaging microscopy. The cytochemical and cytometric aspects of a quantitative ISH procedure were investigated, using human peripheral blood lymphocyte interphase nuclei and probes detecting high copy number target sequences as a model system. These chromosome-specific probes were labeled with biotin, digoxigenin, or fluorescein. The instrumentation requirements are evaluated.

Quantification of the fluorescence ISH signals was performed using an epi-fluorescence microscope with a multi-wavelength illuminator, equipped with a cooled charge couple device (CCD) camera. The performance of the system was evaluated using fluorescing beads and a homogeneously fluorescing specimen.

Specific image analysis programs were developed for the automated segmentation and analysis of the images provided by ISH. Non-uniform background fluorescence of the nuclei introduces problems in the image analysis segmentation procedures. Different procedures were tested. Up to 95% of the hybridization signals could be correctly segmented using digital filtering techniques (min-max filter) to estimate local background intensities.

The choice of the objective lens used for the collection of images was found to be extremely important. High magnification objectives with high numerical aperture, which are frequently used for visualization of fluorescence, are not optimal, since they do not have a sufficient depth of field.

The system described was used for quantification of ISH signals and allowed accurate measurement of fluorescence spot intensities, as well as of fluorescence ratios obtained with double-labeled probes.

CC CYTOLOGY & HISTOLOGY; BIOMETHODS

ST Author Keywords: QUANTIFICATION; CCD CAMERA; IMAGE ANALYSIS; CHROMOSOME POLYMORPHISM

STP KeyWords Plus (R): STAGE ABSORBANCE CYTOPHOTOMETRY; OPTICAL ERRORS; MICROSCOPY; NUCLEI; GLARE

RE Referenced Author (RAU)	Year VOL) (RPG)	
AGARD D A AIKENS R S		353 291	METHOD CELL BIOL

ARNDTJOVIN D J	1985 230	247	SCIENCE
BARROWS G H	984 32	1741	J HISTOCHEM C OCHEM
BAUMAN J	989	1275	FLOW CYTOGENE CS
BENSON D M	1985 100	1309	J CELL BIOL
DUIJNDAM W A L	1980 28	388	J HISTOCHEM CYTOCHEM
DUIJNDAM W A L	1980 28	395	J HISTOCHEM CYTOCHEM
FRANCON M	1961	1	PROGR MICROSCOPY
HIRAOKA Y	1987 238	36	SCIENCE
INOUE S	1986		VIDEO MICROSCOPY
JOHNSON G D	1982 55	231	J IMMUNOL METHODS
JOVIN T M	1989 18	271	ANNU REV BIOPHYS BIO
MAYALL B H	1970	171	INTRO QUANTITATIVE C
NEDERLOF P M	1992 13		CYTOMETRY
NEDERLOF P M	1992 13		CYTOMETRY
NUNEZ D J	1989 263	121	BIOCHEM J
RIDLER T W	1978 8	630	IEEE T SYST MAN CYB
SMITH L C	1986 129	857	METHOD ENZYMOL
TANKE H J	1980 28	11007	J HISTOCHEM CYTOCHEM
TRASK B	1988 78	251	HUM GENET
VANDEKKEN H	1990 11	153	CYTOMETRY
VERBEEK P W	1988 15	1249	SIGNAL PROCESS

L30 ANSWER 16 OF 16 SCISEARCH COPYRIGHT 2000 ISI (R)

AN 92:500902 SCISEARCH

GA The Genuine Article (R) Number: JJ759

TI BEHAVIOR OF PERIOD-ALTERED CIRCADIAN-RHYTHM MUTANTS OF DROSOPHILA IN LIGHT

- DARK CYCLES (DIPTERA, DROSOPHILIDAE)

AU HAMBLENCOYLE M J; WHEELER D A; RUTILA J E; ROSBASH M; HALL J C (Reprint)

CS BRANDEIS UNIV, DEPT BIOL, 235 BASSINE BLDG, WALTHAM, MA, 02254

CYA USA

SO JOURNAL OF INSECT BEHAVIOR, (JUL 1992) Vol. 5, No. 4, pp. 417-446. ISSN: 0892-7553.

DT Article; Journal

FS AGRI

LA ENGLISH

REC No References

Keyed

Adults of Drosophila melanogaster had their locomotor activity monitored under conditions of cycling light and dark (12 h each per cycle). The elementary behavior of wild-type flies under these "LD" conditions fluctuated between levels of high and levels of low activity. Two high-activity peaks occurred within a given cycle: one at about dawn; the other, at around dusk. Such accentuated activity levels gradually subsided to troughs in the middle of the day and of the night, after

which

the flies anticipated the next environmental transition by gradually becoming more active. Descriptions of these activity profiles were augmented by newly developed formal analyses of the "diel rhythm" phases (based in part on digital filterings of the raw behavioral data). The applications of these analyses led to objective, automated determination of when in the morning and the evening the flies' activity peaks occur. This normal diel behavior was compared to the locomotor activity and phase determinations for a series of rhythm variants. Most of these involved mutations at the period (per) locus and germ-line transformants bearing normal or altered forms of DNA cloned from this "clock gene." Such genetic variants have been shown previously to exhibit, in constant darkness, strain-specific circadian periods ranging from about 19 to about 29 h. We now show that the phases of the evening peaks of activity under LD conditions were correspondingly earlier than normal for the short-period mutants and later than normal

for

those with long circadian cycle durations. The morning peaks, however, moved (in comparison to the normal phase position) minimally under the influence of a given per variant.

CC ENTOMOLOGY

Author Keywords: I COMOTOR ACTIVITY; PER-SHORT MUTA; PER-LONG MUTANTS; PER-TRANSGENICS; CK MUTANT; BLIND NORPA MUTANT; ASE ANALYSIS

35 ANSWER 1 OF 3 BIOSIS COPYRIGHT 2000 BIOSIS DUPLICATE 1 1996:541008 BIOSIS AN PREV199699263364 DN Steady-state analysis of somatosensory evoked potentials. ΤI Noss, Roger S. (1); Boles, Colby D.; Yingling, Charles D. AU (1) Dep. Anesthesia, Sch. Med., Univ. California, San Francisco, CA CS 94143-0648 USA Electroencephalography and Clinical Neurophysiology, (1996) Vol. 100, No. SO 5, pp. 453-461. ISSN: 0013-4694. Article DTEnglish LAWe report the development of a new method for frequency domain analysis AB of steady-state somatosensory evoked potentials (SEPs) to amplitude-modulated electrical stimulation, which can be recorded in significantly less time than traditional SEPs. Resampling techniques were used to compare the steady-state SEP to traditional SEP recordings, which are based on signal averaging in the time domain of cortical responses to repetitive transient stimulation and take 1-2 min or more to obtain a satisfactory signal/noise ratio. Median nerves of 3 subjects were stimulated continuously with electrical alternating current at several modulation frequencies from 7 to 41 Hz. Amplitude modulation was used to concentrate the power in higher frequencies, away from the modulation frequency, to reduce the amount of stimulus artifact recorded. Data were tested for signal detectability in the frequency domain using the T-circ-2 statistic. A reliable steady-state response can be recorded from scalp electrodes overlying somatosensory cortex in only a few seconds. In contrast, no signal was statistically discriminable from noise in the transient SEP from as much as 20 s of data. This dramatic time savings accompanying steady-state somatosensory stimulation may prove useful for monitoring in the operating room or intensive care unit. Biophysics - General Biophysical Techniques CC Nervous System - General; Methods *20501 Nervous System - Physiology and Biochemistry *20504 Hominidae *86215 ВC Major Concepts ΙT Methods and Techniques; Nervous System (Neural Coordination) Miscellaneous Descriptors ΙT ANALYTICAL METHOD; NERVOUS SYSTEM; NEW METHOD; SIGNAL DETECTABILITY; SOMATOSENSORY EVOKED POTENTIAL; STEADY-STATE ANALYSIS ORGN Super Taxa Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name human (Hominidae) ORGN Organism Superterms animals; chordates; humans; mammals; primates; vertebrates L35 ANSWER 2 OF 3 BIOSIS COPYRIGHT 2000 BIOSIS DUPLICATE 2 1991:384339 BIOSIS ANDN BA92:61654 PITUITARY MICROCIRCULATION PHYSIOLOGICAL ASPECTS AND CLINICAL IMPLICATIONS A LASER-DOPPLER FLOW STUDY DURING TRANSSPHENOIDAL ADENOMECTOMY. STEINMEIER R; FAHLBUSCH K; POWERS A D; DOETTERL A; BUCHFELDER M ΑU

CS NEUROCHIRURGISCHE KLINK DER UNIV. ERLANGEN-NUERNBERG, SCHWABACHANLAGE 6 KOPFKLINIKUM, 852 ERLANGEN, GERMANY.

SO NEUROSURGERY (BALTORE), (1991) 29 (1), 47-54. CODEN: NRSRDY.

FS BA; OLD

LA English

The anterior and posterior putuitary lobes (AL and PL, respectively) are AB assumed to differ in the type of vacular supply and structure of their microvascular networks. Animal experiments have shown that the pituitary microvascular flow differs between the two lobes, being extremely high in the PL and low in the AL. For technical reasons, it has hitherto not been possible to study pituitary microflow in humans. Laser-Doppler flowmetry (LDF) is now a well-established method for real-time monitoring of microcirculation, applicable also in humans. In a prospective clinical study, the microflow in the AL and PL was measured during transsphenoidal microsurgery in 52 patients with adenomas of different size, growth characteristics, and endocrinological activity. The mean microflow in the PL (177.7 .+-. 12.6 [flux]) was found to be about six times higher than that in AL (27.4 .+-. 2.7 [flux]). No difference in the laser-Doppler fractional volume of the lobes could be detected (0.73 .+-. 0.06 [] vs. $0.77 \cdot +- \cdot \cdot \cdot 0.07$ [], where [] designates the ratio of the alternating current output to the direct current output signals). Microflow within the pituitary lobes was influenced neither by the histological type nor the size of the adenoma. Additionally, LDF signal-averaging triggered by the electrocardiogram allowed detection of different characteristic pulsatile microvascular

flow

patterns in the AL and PL. Our findings provide strong physiological support for the idea that the angioarchitecture of the pituitary lobes differs. With this method, the AL and PL can be identified objectively during surgery. LDF might provide useful information concerning intraoperative surgical approach.

CC Radiation - Radiation and Isotope Techniques *06504
Biophysics - General Biophysical Techniques *10504
Anatomy and Histology, General and Comparative - Surgery *11105
Anatomy and Histology, General and Comparative - Radiologic Anatomy *11106

Anatomy and Histology, General and Comparative - Microscopic and Ultramicroscopic Anatomy 11108

Cardiovascular System - Physiology and Biochemistry *14504 Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies *15002

Endocrine System - Pituitary *17014

Neoplasms and Neoplastic Agents - Pathology; Clinical Aspects; Systemic Effects *24004

Neoplasms and Neoplastic Agents - Therapeutic Agents; Therapy *24008

BC Hominidae 86215

IT Miscellaneous Descriptors

HUMAN VASCULAR SUPPLY ANGIOARCHITECTURE

- L35 ANSWER 3 OF 3 EMBASE COPYRIGHT 2000 ELSEVIER SCI. B.V.
- AN 85095295 EMBASE
- DN 1985095295
- TI Highly sensitive microcomputer-controlled ac magnetometer with a phase locked data acquisition system.
- AU Martin W.E.; Wieser J.
- CS Sektion Physik, University of Munich, Munich, Germany
- SO Journal of Physics E: Scientific Instruments, (1985) 18/4 (342-349). CODEN: JPSIAE
- CY United Kingdom
- DT Journal
- FS 027 Biophysics, Bioengineering and Medical Instrumentation
- LA English
- AB A highly sensitive microcomputer-controlled magnetometer for AC measurements in applied fields of up to 3 x 105 A m-1 is described. The

'AC magnetometer' (operating frequency 50 Hz nominally) is based on a mains-powered sole id, and a high resolution signa averaging system detable of analysing extremely small signals down to the order of electronic noise (about 1 .mu.V). The high density of data points allowed by the system demands a personal computer acting as control unit for data acquisition (based on linear summation averaging) and data handling. To profit by the sensitivity and resolution capacity given by signal averaging methods and to guarantee precise operation of the mains-supplied AC magnetometer, the data acquisition process must be exactly synchronised to power line frequency. In order to meet this basic requirement in spite of random line frequency fluctuations up to .+-.2.per thousand., a line locked oscillator circuit acting as averager system clock has been developed. The circuit is described here in detail for the first time. The AC magnetometer was employed to record the magnetisation curves, M against H, of ferromagnetic samples having small magnetically effective cross sections, and to determine their AC magnetic properties (saturation magnetisation, remanence, coercivity, susceptibility) in the temperature range down to 4.2 K. The performance of the system is demonstrated here by some tests and by presenting results οf magnetic measurements showing e.g. the interesting magnetic behaviour of highly concentrated metal-hydrogen systems with the ferromagnetic component nickel (here with magnetic cross sections down to about 5 x 10-5 mm2). Medical Descriptors: CT*alternating current *data analysis *magnetometer *microcomputer computer analysis

36 ANSWER 1 OF 1 EMBASE COPYRIGHT 2000 ELSEVIER SCI. B.V. 1999199911 EMBASE ΑN Some design concepts for electrical impedance measurement. TIGoovaerts H.G.; Faes Th.J.C.; Raaijmakers E.; Heethaar R.M. ΑU H.G. Goovaerts, Dept. Clinical Physics Informatics, Inst. Cardiovascular CS Research ICarVU, Univ. Hospital Vrije Universiteit, 1007 MB Amsterdam, Netherlands Annals of the New York Academy of Sciences, (1999) 873/- (388-395). SO Refs: 7 ISSN: 0077-8923 CODEN: ANYAA United States CY Journal; Conference Article \mathtt{DT} Radiology FS 014 Biophysics, Bioengineering and Medical Instrumentation 027 English LAEnglish \mathtt{SL} Design concepts for the implementation of two basic functions for AB measurement of electrical impedance are presented: current injection and voltage measurement. At relatively high frequencies, the application of an alternating current through the body or a body segment results in electromagnetic stray fields that reduce the amount of current actually injected into the tissue under study. It is shown that electrical isolation and small dimensions of the isolated section are indispensable in order to substantially reduce these stray currents. The paper describes a new wideband current source configuration driven by direct digital sine wave synthesis (DDS) presenting very low stray currents due to a symmetrical layout. Two implementations of the actual current source circuit are presented: (1) a voltage-controlled system and (2) a current conveyor-based circuit. A wideband input amplifier with transformer coupling is described. The current source, amplifier, and (in case of tomography) multiplexer are also situated on an electrically isolated front end. The presented concepts are applied in a new electrical impedance tomograph (EIT) presently under construction in our department. Medical Descriptors: CT*impedance *diagnostic imaging alternating current electric potential measurement electromagnetic field direct current digital filtering amplifier tomography human confe

DUPLICATE 1 L41 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2000 ACS 1996:554604 CAPLUS AN 125:259590 DN Surface charge density measurements with a controlled growth mercury TIelectrode O'Dea, John J.; Ciszkowska, Malgorzata; Osteryoung, Robert A. ΑU Dep. Chem., North Carolina State Univ., Raleigh, NC, 27695, USA CS Electroanalysis (1996), 8(8-9), 742-747 SO CODEN: ELANEU; ISSN: 1040-0397 Journal DT English LA72-2 (Electrochemistry) CC Section cross-reference(s): 66 The surface charge d. of the Hg electrolyte interface is estd. by using AΒ chronocoulometry at a controlled-growth Hg electrode. After initial formation and equilibration, the Hg drop is expanded by further addn. of Hq. Direct measurement of the charge, required as new area is formed, is used to est. the surface charge d. The Hg drop is modeled as a step-wise expanding sphere. The capillary noise continuously produced by stationary drops under potential control was investigated and characterized. Spectral anal. of the noise reveals that the electrode is particularly sensitive to vibrations near the resonant frequency of the suspended drop. Ambient vibrations in the lab. environment produce alternating currents at the electrode which vanish at the point of zero charge and so mark its position. surface charge density mercury electrode STElectrodes IT(surface charge d. detn. of Hg/electrolyte interface by chronocoulometry at a controlled-growth Hg electrode) Electric charge IT (surface, d.; detn. of Hg/electrolyte interface by chronocoulometry at a controlled-growth Hg electrode) 7601-90-3, Perchloric acid, uses ITRL: NUU (Nonbiological use, unclassified); PRP (Properties); USES (Uses) (detn. of Hg/HClO4 electrolyte interface by chronocoulometry at controlled-growth Hg electrode) 7439-97-6, Mercury, uses ITRL: DEV (Device component use); PRP (Properties); USES (Uses) (surface charge d. detn. of Hg/electrolyte interface by chronocoulometry at a controlled-growth Hg electrode) L41 ANSWER 2 OF 10 CAPLUS COPYRIGHT 2000 ACS 1991:16654 CAPLUS AN114:16654 DN Spectral analysis using a high-voltage a.c. arc. TIStability of the constant calibration diagram Skuratova, T. A.; Trapitsyn, N. F. ΑU Kirg. Gos. Univ., Frunze, USSR CS Zh. Prikl. Spektrosk. (1990), 53(4), 662-3 SO CODEN: ZPSBAX; ISSN: 0514-7506 Journal DTRussian LA79-1 (Inorganic Analytical Chemistry) CC A high-voltage a.c. arc source was used in the anal. of brass LS-59 for AB impurities with the utilization of a const. calibration diagram. The method of carrying out the analyses was developed in an earlier work.

The

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const. calibration diagram was constructed with std. samples contg. Si,
    Sn, Fe, Pb, Al, an Ni, which were subjected to committion in the arc at
a
    current of 3 A and source voltage of 1900 V, with a distance between the
    electrodes of 3 mm. The std. sample served as the lower
     electrode, while the upper electrode was C. With this
    method, it is possible to keep position of this diagram const. over a
long
     time. In 16 yr of continuous operation of the generator, no displacement
     of the calibration diagram was obsd. This source for the excitation of
     spectra can be used idefinitely for the anal. of metals and alloys.
     high voltage alternating current arc analysis; metal
ST
     analysis alternating current arc; alloy analysis
     alternating current arc; const calibration diagram
     stability analysis
    Alloys, analysis
IT
     Metals, analysis
     RL: ANT (Analyte); ANST (Analytical study)
        (anal. of, stability of const. calibration diagram using high-voltage
        a.c. arc source for)
     Spectrochemical analysis
IT
        (at. emission, of alloys and metals, stability of const. calibration
        diagram using high-voltage a.c. arc source for)
L41 ANSWER 3 OF 10 CAPLUS COPYRIGHT 2000 ACS
    1983:532822 CAPLUS
AN
     99:132822
DN
     Characteristics of an alternating-current arc and
ΤI
     high-frequency spark discharges used in the spectral
     analysis of metals and alloys in air and argon
     Eroshenko, L. E.; Dem'yanchuk, A. S.
ΑU
CS
     USSR
     Zh. Prikl. Spektrosk. (1983), 39(1), 15-21
SO
     CODEN: ZPSBAX; ISSN: 0514-7506
     Journal
\mathsf{DT}
     Russian
LA
     79-2 (Inorganic Analytical Chemistry)
CC
     Discharge generated with conical Cu counter electrodes in anal.
AΒ
     of steel and cast iron stds. were studied by high-speed filming and also
     by metallog. and recording profiles of the analyzed samples. Suppression
     of matrix effects with a.c. arc sources in Ar and high-frequency spark
     sources in air and Ar can be explained by random movement of the
discharge
     on the sample surface.
     arc discharge metal analysis spectrog; spark discharge metal analysis
ST
     spectrog; metal analysis spectrog discharge source; alloy analysis
     spectrog discharge source; air atm spectrog discharge; argon atm spectrog
     discharge
     Alloys, analysis
IT
     Metals, analysis
     RL: ANT (Analyte); ANST (Analytical study)
         (anal. of, matrix effect suppression with arc and spark sources in
         spectroq.)
     Electric arc
 IT
     Electric spark
         (in spectrog. anal. of alloys and metals, in air and argon, matrix
         effect suppression with)
      Spectrochemical analysis
 IT
         (emission, of alloys and metals, matrix effect suppression with arc
 and
         spark sources in)
                             12597-69-2, analysis
      11097-15-7, analysis
 IT
      RL: ANT (Analyte); ANST (Analytical study)
         (anal. of, matrix effect suppression with arc and spark sources in
         spectrog.)
```

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L41 ANSWER 4 OF 10 CALUS COPYRIGHT 2000 ACS
     1984:603338 CAPLU
AN
     101:203338
DN
    Vaporization of basic components of a powdered sample from an
TI
     alternating-current carbon arc crater
     Yankovskaya, T. A.
ΑU
     USSR
CS
     Primen. Spektr. Anal. Nar. Khoz. Nauchn. Issled., Mater. Resp. Semin.
SO
     Spektr. Anal. (1982), Meeting Date 1981, 72-8. Editor(s): Petukh, M. L.;
     Yankovskii, A.. Publisher: Akad. Nauk BSSR, Inst. Fiz., Minsk, USSR.
     CODEN: 52GVA2
     Conference
DT
LA Russian
    79-1 (Inorganic Analytical Chemistry)
CC
     The erosion of arc electrodes and the evapn. of sample
AB
     components in emission spectrog. anal. of powders with 9 and 18-A sources
     were studied as functions of time by using electrodes of
     different dimensions for anal. of model mixts. of synthetic silicate rock
     stds. of different wt.
     sample evapn arc spectral analysis; powder evapn arc
ST
     spectral analysis; carbon arc powd sample evapn;
     silicate rock analysis emission spectrometry
     Powders
\operatorname{IT}
        (anal. of, by emission spectrometry, vaporization of basic components
        in a.c. arcs for)
     Erosion
IT
        (of arc electrodes in spectrochem. anal.)
     Evaporation
IT
         (of basic components of powd. samples from a.c. arcs for
      spectral anal.)
     Electric lamps
IT
         (arc, with carbon electrodes, for emission spectral
      anal., erosion and vaporization studies in relation to)
     Spectrochemical analysis
IT
     Spectrochemical analysis
         (emission, of powd. samples, vaporization in a.c. arc sources in
        relation to)
     Rocks
IT
     RL: ANT (Analyte); ANST (Analytical study)
         (silicate, anal. of, by emission spectrometry, vaporization of basic
        components in a.c. arcs for)
     7440-44-0, uses and miscellaneous
IT
     RL: USES (Uses)
         (arc electrodes, in emission spectral anal
         ., vaporization of basic components of powd. samples from a.c.)
     ANSWER 5 OF 10 CAPLUS COPYRIGHT 2000 ACS
L41
      1978:452689 CAPLUS
AN
     89:52689
DN
      Study of the relation of effective parameters of an alternating-
\mathtt{TI}
      current arc plasma to the composition of coal graphite materials
      Zhilova, A. N.; Akimov, V. A.; Egorova, V. A.
ΑU
      USSR
 CS
      Tr. Mosk. Khim.-Tekhnol. Inst. (1976), 91, 129-30
 SO
      CODEN: TMKIAT; ISSN: 0371-9723
      Journal
 DT
      Russian
 LA
      79-1 (Inorganic Analytical Chemistry)
 CC
      Section cross-reference(s): 76
      The effective parameters of an 8-A a.c. arc plasma between graphite
 AB
      electrodes for emission spectral anal. were
      calcd. as functions of the analyte concns. in carbonaceous samples, such
      as coke, glassy C, graphite, and pyrolytic graphite. The effective temp.
      (Te) was calcd. by using Zn as a thermometric element, the presence of
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which was maintained in the plasma by using a 2-mm wide and 15-mm deep electrode crater. The electron concn. (ne) was call by using the spectral line tensities of Mg. Both Te and nedecrease symbatically with an increase in analyte concn. from 3 .times. 10-3 to 1 .times. 10-1%. The type of coke used did not affect ne and Te at analyte concns. .ltoreq.2 .times. 10-1%. carbon analysis emission spectrometry; coke analysis emission stspectrometry; graphite analysis emission spectrometry; temp spectral arc plasma; electron concn spectral arc plasma; arc plasma temp electron concn Coke ITRL: ANT (Analyte); ANST (Analytical study) (anal. of, by emission spectrometry, electron concn. and plasma temp. in a.c. arc for) Plasma IT(electron concn. and temp. of arc, analyte concn. in spectrochem. anal. of carbonaceous material in relation to) Spectrochemical analysis ΙT (emission, of carbonaceous materials, electron concn. and plasma temp. in a.c. arc for) 7440-44-0, analysis ITRL: ANST (Analytical study) (anal. of glassy, by emission spectrometry, electron concn. and plasma temp. in a.c. arc for) 7782-42-5, analysis ΙT RL: ANT (Analyte); ANST (Analytical study) (anal. of, by emission spectrometry, electron concn. and plasma temp. in a.c. arc for) L41 ANSWER 6 OF 10 CAPLUS COPYRIGHT 2000 ACS 1978:452679 CAPLUS AN89:52679 DN Axial-time distribution of the basic plasma parameters in an ΤI alternating current arc. II Kapitonov, A. N. ΑU USSR CS Nek. Vopr. Fiz. (1975), 9-14. Editor(s): Solov'ev, G. N. Publisher: SO Yakutsk. Gos. Univ., Yakutsk, USSR. CODEN: 3700AY Conference DT LARussian 79-1 (Inorganic Analytical Chemistry) CCThe axial and time distribution of the electron concn. (Ne) and temp. (T) ABin the plasma of an arc for emission spectral anal. is discussed. The excitation conditions characterized by Ne and T depend on the phase of the arc discharge, electrode polarity, and the position in the arc gap. Stable excitation conditions were obsd. at 0.8-1.7 mm above the lower electrode. arc plasma emission spectral analysis; temp \mathtt{ST} distribution arc plasma; electron distribution arc plasma Plasma ${ t IT}$ (electron concn. and temp. in a.c. arc, axial-time distribution of) Spectrochemical analysis IT (emission, excitation condition in a.c. arc for) ANSWER 7 OF 10 CAPLUS COPYRIGHT 2000 ACS L411978:452678 CAPLUS AN89:52678 DN Axial-time distribution of the plasma parameters in an alternating TIcurrent arc. I Alekseev, M. A.; Kapitonov, A. N. ΑU

USSR

CS

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Nek. Vopr. Fiz. (1975), 3-8. Editor(s): Solov'ev, G. N. Publisher:
SO
                         Yakutsk, USSR.
    Yakutsk. Gos. Univ
    CODEN: 3700AY
     Conference
DT
     Russian
LA
    79-1 (Inorganic Analytical Chemistry)
CC
     The axial and time distribution of the electron concn. (Ne) and temp. (T)
AB
     in the plasma of an a.c. arc for emission spectral anal
     . was explained by the effect of the elec. field upon the transport of
     metal atoms in the excitation zone. The distribution of Ne and T is a
     function of the polarity of the graphite electrode contg. the
     sample. The excitation conditions during the anal. of samples contg.
     easily ionizable components, such as Na or Ca, can be improved by placing
     the easily ionizable materials on both electrodes.
     arc plasma emission spectral analysis; temp
ST
     distribution arc plasma; electron distribution arc plasma
     Plasma
IT
        (electron concn. and temp. in a.c. arc, transport of metals in elec.
        field in relation to)
     Electric field, chemical and physical effects
IT
        (alternating, on transport of metals in arc plasma, axial and time
        distribution of plasma parameters in relation to)
     Spectrochemical analysis
IT
         (emission, for easily ionizable components, a.c. arc plasma
        stabilization by placing materials on both electrodes in)
L41 ANSWER 8 OF 10 CAPLUS COPYRIGHT 2000 ACS
     1972:549675 CAPLUS
AN
     77:149675
DN
     Spectral analysis of plant material
TI
     Glinski, Jan; Nowicki, Ryszard
ΑU
     Inst. Agrophys., Pol. Acad. Sci., Lublin, Pol.
CS
     Pol. J. Soil Sci. (1972), 4(2), 113-18
SO
      CODEN: PJSOBN
     Journal
\mathsf{DT}
     English
{
m LA}
     11-1 (Plant Biochemistry)
CC
      Section cross-reference(s): 9
     Thirteen trace elements (B, Ba, Co, Cu, Cr, F, Mn, Mo, Ni, Pb, Sr, V, and
AB
      Zn) in plant ash can be detd. by emission spectral anal
      . with a spectrograph of medium dispersion Q-24 on sample excitation from
      shallo craters of graphite electrodes in an interrupted arc of
      alternating current, with synthetic stds. Six major
      elements (Al, K, Mg, Na, P, and Si) can be detd. simultaneously.
      trace element detn plant; mineral detn plant
 ST
      Trace elements
 IT
      RL: ANT (Analyte); ANST (Analytical study)
         (detn. of, in plant tissue)
      Plant analysis
 {	t IT}
         (for trace elements)
      ANSWER 9 OF 10 CAPLUS COPYRIGHT 2000 ACS
 L41
      1969:117717 CAPLUS
 AN
      70:117717
 DN
      Spectrographic study in the ceramic industry
 ΤI
      Bolgar, Gabor
 ΑU
      Magnezitipari Muevek, Budapest, Hung.
 CS
      Kohasz. Lapok (1967), 100(5), 217-8
 SO
      From: CZ 1969 (3), Abstr. No. 1983
      CODEN: KOLAAR
      Journal
 \mathtt{DT}
      Hungarian
 \mathtt{L}\!\mathtt{A}
      57 (Ceramics)
 CC
      The powd. sample is mixed in an agate mortar with Ba(NO3)2 and spectrally
 AΒ
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pure graphite powder. The absorption is measured with a graphite

electrode and alternating current arc The las Mg 2781.42 A. (for Si, Mn ar Fe) and Ba 3071.59 excitation. Α. (for Al and Ca) are used as standards for comparison. The process gives an acceptable value at high-Mg content. spectral anal ceramics STCeramic materials IT(anal. of, spectrochem.) L41 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2000 ACS 1967:34491 CAPLUS AN66:34491 DN Effect of electrode polarity on the current and radiation of an TI alternating current arc for spectral analysis Brainin, E. I.; Pyasetskaya, L. I. AU Zh. Prikl. Spektrosk. (1966), 5(3), 399-402 SO CODEN: ZPSBAX Journal $\mathsf{D}\mathbf{T}$ LARussian 79 (Inorganic Analytical Chemistry) CC The relation between spectral line intensities and the current of an AB a.-c. arc during consecutive half cycles was studied exptl. by using the ISP-30 spectrograph with electrodes of pure C and Ag, and Ag electrodes contq. various amts. of CdO, Al2O3, ZnO, Fe2O3, CuO, Na2CO3, Cu, Ni, and W. The current difference (.DELTA.i) and the log of spectral line intensities were detd. as a function of temp. and the admixt. concns. for 20 electrodes during the anode-cathode alternation. The difference in .DELTA.i during the alternation is attributed to the difference in the emission of electrons and ions from the electrodes into the plasma. The emission of electrons increased with cathode temp. When both electrodes are of the same material, then the difference in the emission is detd. by the electrode surface temp. In most cases the spectra intensities increased with increasing current. ELECTRODES POLARITY SPECTROSCOPY; POLARITY ELECTRODES \mathtt{ST} SPECTROSCOPY; SPECTROSCOPY ELECTRODES POLARITY Analysis IT

(spectrochem., electrode polarity effect on current and

radiation of a.c. arc in)